Connecting via Winsock to STN

Welcome to STN International! Enter x:X

LOGINID:SSPTASYG1600

PASSWORD:

PASSW0 TERMIN		ENTE	ER 1	, 2, 3, OR ?):2					
* * *	* *	* *	* *	* Welcome to STN International * * * * * * * * *					
NEWS	1			Web Page for STN Seminar Schedule - N. America					
NEWS	2	JAN	02	STN pricing information for 2008 now available					
NEWS	3	JAN	16	CAS patent coverage enhanced to include exemplified					
				prophetic substances					
NEWS	4	JAN	28	USPATFULL, USPAT2, and USPATOLD enhanced with new					
				custom IPC display formats					
NEWS		JAN		MARPAT searching enhanced					
NEWS	6	JAN	28	USGENE now provides USPTO sequence data within 3 days of publication					
NEWS	7	JAN.	28	TOXCENTER enhanced with reloaded MEDLINE segment					
NEWS				MEDLINE and LMEDLINE reloaded with enhancements					
NEWS				STN Express, Version 8.3, now available					
				PCI now available as a replacement to DPCI					
NEWS				IFIREF reloaded with enhancements					
NEWS				IMSPRODUCT reloaded with enhancements					
NEWS				WPINDEX/WPIDS/WPIX enhanced with ECLA and current					
Hand	13			U.S. National Patent Classification					
NEWS	14	MAR	31	IFICDB, IFIPAT, and IFIUDB enhanced with new custom					
				IPC display formats					
NEWS	15	MAR	31	CAS REGISTRY enhanced with additional experimental					
				spectra					
NEWS	16	MAR	31	CA/CAplus and CASREACT patent number format for U.S. applications updated					
NEWS	17	MAR	21	LPCI now available as a replacement to LDPCI					
NEWS				EMBASE, EMBAL, and LEMBASE reloaded with enhancements					
NEWS				STN AnaVist, Version 1, to be discontinued					
NEWS		APR							
NEWS	20	APK	13	15 WPIDS, WPINDEX, and WPIX enhanced with new predefined hit display formats					
NEWS	21	APR	28	EMBASE Controlled Term thesaurus enhanced					
NEWS				IMSRESEARCH reloaded with enhancements					
NEWS				INPAFAMDB now available on STN for patent family					
112110	23		50	searching					
NEWS	24	MAY	3.0	DGENE, PCTGEN, and USGENE enhanced with new homology					
				sequence search option					
NEWS	25	JUN	06	EPFULL enhanced with 260,000 English abstracts					
NEWS		JUN		KOREAPAT updated with 41,000 documents					
				, , , , , , , , , , , , , , , , , , , ,					
NEWS	EXPRESS			RUARY 08 CURRENT WINDOWS VERSION IS V8.3,					
			AND	CURRENT DISCOVER FILE IS DATED 20 FEBRUARY 2008					
NEWS	HOURS		ST	N Operating Hours Plus Help Desk Availability					
	LOGIN			Welcome Banner and News Items					
	S IPC8		For general information regarding STN implementation of IPC 8						
				3					

Enter NEWS followed by the item number or name to see news on that specific topic.

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* * * * * * * * * * * * * * * * STN Columbus * * * * * * * * * * * * * * * * * *

FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008

=> b hcap

COST IN U.S. DOLLARS

SINCE FILE TOTAL.

ENTRY SESSION FULL ESTIMATED COST 0.42 0.42

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 12 Jun 2008 VOL 148 ISS 24 FILE LAST UPDATED: 11 Jun 2008 (20080611/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> us20050249857/pn

US20050249857 IS NOT A RECOGNIZED COMMAND

The previous command name entered was not recognized by the system. For a list of commands available to you in the current file, enter "HELP COMMANDS" at an arrow prompt (=>).

=> set autosearch on SET COMMAND COMPLETED

=> us20050249857/pn

1 US20050249857/PN L1

=> b reg;tra rn COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION FILL ESTIMATED COST 2.69 3.11

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 11 JUN 2008 HIGHEST RN 1027436-61-8 DICTIONARY FILE UPDATES: 11 JUN 2008 HIGHEST RN 1027436-61-8

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 9, 2008.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

```
L2 TRANSFER L1 1- RN: 16 TERMS
L3 16 L2

=> 13 and >= 2 46.150.18/rid
16596122 RID.CNT >= 2
22121069 46.150.18/RID
11101089 >= 2 46.150.18/RID
(RID.CNT >= 2 (T) 46.150.18/RID)
L4 10 L3 AND >= 2 46.150.18/RID
```

=> d sca

14 10 ANSMERS REGISTRY COPYRIGHT 2008 ACS on STN Phenol, 4,4"-(tetrahydro-1H,3H-furo[3,4-c]furan-1,4-diyl)bis[2,6-dimethoxy-,(1R,3a5,4R,6a5)-relpt C22 H26 08

Relative stereochemistry.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> 13 and oc4-c6-c6/es 58515 OC4-C6-C6/ES

1 L3 AND OC4-C6-C6/ES

=> d sca

L5

L5 1 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Naphtho[2,3-c]furan-1(3H)-one, 3a,4,9,9a-tetrahydro-6-hydroxy-4-(4-hydroxy-3-methoxyphenyl)-7-methoxy-, (3aR,4S,9aR)-

MF C20 H20 O6

Absolute stereochemistry. Rotation (-).

^{**}PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT**

ALL ANSWERS HAVE BEEN SCANNED

=> 13 and 6-c6/es 0 6-C6/ES

L6 0 L3 AND 6-C6/ES

=> del 16 DELETE L6? (Y)/N:v

=> 13 and c6-c6/es

1122454 C6-C6/ES L6 1 L3 AND C6-C6/ES

=> d sca

L6 1 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN 2,3-Naphthalenedimethanol, 1,2,3,4-tetrahydro-7-hydroxy-1-(4-hydroxy-3-methoxyphenyl)-6-methoxy-, (1S,2R,3R)-

MF C20 H24 O6

CI COM

Absolute stereochemistry. Rotation (+).

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

ALL ANSWERS HAVE BEEN SCANNED

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

L3 L4 16 SEA L2 10 L3 AND >=2 46.150.18/RID L5 1 L3 AND OC4-C6-C6/ES L6 1 L3 AND C6-C6/ES => str :gra c3 :dis C-~ C-~ C :nod 1 3 cb, dis Cb--^ C-- Cb :nod 2 gl,gra c3,c1,c2,c1,arr,dis C 7 C~~C C 10 Cb\G1\Cb Cb C\C\C C C :nod 4 6 7 9 ak, 5 8 10 hy :dis sia $\texttt{Cb} \backsim \texttt{G1} \backsim \texttt{Cb} \qquad \texttt{Ak} \backsim \texttt{Hy} \backsim \texttt{Ak} \qquad \qquad \texttt{Ak} \enspace 7 \qquad \texttt{Hy} \backsim \texttt{Ak}$ Hy 10 4 5 6 ลี 9 1 2 3 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 10 STEREO ATTRIBUTES: NONE :eco 5 8 el o e4 c,dis sia Ak 7 Hy~ Ak Cb~G1~Cb Ak~Hy~Ak Hy 10 4 5 6 1 2 3 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED ECOUNT IS E4 C E1 O AT 5 ECOUNT IS E4 C E1 O AT 8 GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 10 STEREO ATTRIBUTES: NONE :eco 7 m3-x5 c,dis sia Ak 7 Hy~ Ak Hy 10 Cb√G1√Cb Ak√Hv√Ak 1 2 3 4 5 6 8 9

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED ECOUNT IS E4 C E1 O AT 5 ECOUNT IS M3-X5 C AT 7 ECOUNT IS E4 C E1 O AT 8

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 10

STEREO ATTRIBUTES: NONE :eco 10 e2 o e6 c,dis sia

NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT BCLEVEL IS LIMITED
ECOUNT IS E4 C E1 0 AT 5
ECOUNT IS M3-X5 C AT 7
ECOUNT IS E4 C E1 0 AT 8
ECOUNT IS E6 C E2 0 AT 10

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 10

STEREO ATTRIBUTES: NONE :ggc 10 pcy :dis sia

NODE ATTRIBUTES:

DEFAULT MLEVEL IS ATOM

GGCAT IS PCY AT 10

DEFAULT ECLEVEL IS LIMITED

DEFAULT ECLEVEL IS LIMITED

ECOUNT IS 84 C E1 O AT 5

ECOUNT IS 84 C E1 O AT 8

ECOUNT IS 86 C E2 O AT 10

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 10

STEREO ATTRIBUTES: NONE :var g1=4-1 6-3/7/8-1 9-3/10 :dis sia

Cb G1 Cb Ak Hy Ak Ak @7 Hy Ak Hy @10

VAR G1=4-1 6-3/7/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E4 C E1 0 AT 5

```
ECOUNT IS M3-X5 C AT 7
ECOUNT IS E4 C E1 O AT 8
ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 10
STEREO ATTRIBUTES: NONE
:end
L7 STRUCTURE CREATED
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
               TRA L1 1- RN :
                                    16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
            16 SEA L2
            10 L3 AND >=2 46.150.18/RID
L4
L5
             1 L3 AND OC4-C6-C6/ES
L6
            1 L3 AND C6-C6/ES
L7
               STR
=> 17
SAMPLE SEARCH INITIATED 14:36:49 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 1779298 TO ITERATE
 0.1% PROCESSED
                   2000 ITERATIONS
                                                              29 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**
                       BATCH **INCOMPLETE**
PROJECTED ITERATIONS:
                       35522387 TO 35649533
PROJECTED ANSWERS:
                          506385 TO 525607
T.R
            29 SEA SSS SAM L7
=> >=2 46.150.18/rid
      16596122 RID.CNT >= 2
      22121069 46.150.18/RID
SYSTEM LIMITS EXCEEDED - SEARCH ENDED
The search profile you entered was too complex or gave too many
answers. Simplify or subdivide the query and try again. If you have
```

exceeded the answer limit, enter DELETE HISTORY at an arrow prompt (->) to remove all previous answers sets and begin at L1. Use the SAVE command to store any important profiles or answer sets before

=> d his

using DELETE HISTORY.

```
(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
               TRA L1 1- RN :
                                    16 TERMS
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
1.3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
L6
L7
              STR
L8
             29 L7
=> acr 1839
          2189 ACR
            14 ACRS
          2203 ACR
                (ACR OR ACRS)
          2436 1839
            0 ACR 1839
                (ACR(W) 1839)
=> del 19 y
=> scr 1839
     SCREEN CREATED
L9
=> dhis
            0 DHIS
L10
           0 DHIS
=> del 110 v
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
                SET AUTOSEARCH ON
L1
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
                TRA L1 1- RN : 16 TERMS
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L.4
             10 L3 AND >=2 46.150.18/RID
             1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
1.6
1.7
               STR
T. 8
             29 L7
L9
               SCR 1839
```

```
=> 17 and 19
SAMPLE SEARCH INITIATED 14:40:41 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 1400437 TO ITERATE
 0.1% PROCESSED
                    2000 ITERATIONS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**
                         BATCH **INCOMPLETE**
PROJECTED ITERATIONS:
                         27948931 TO 28068549
PROJECTED ANSWERS:
                            397594 TO 414658
T.10
            29 SEA SSS SAM L7 AND L9
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
                SET AUTOSEARCH ON
L1
               1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
                TRA L1 1- RN : 16 TERMS
   FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
              10 L3 AND >=2 46.150.18/RID
L5
              1 L3 AND OC4-C6-C6/ES
L6
              1 L3 AND C6-C6/ES
L7
                STR
L8
            29 L7
L9
               SCR 1839
L10
            29 L7 AND L9
=> str 17
:dis
                                Ak @7
                                                      Hy @10
Cb~G1~Cb Ak~Hy~Ak
                                          Hy∽ Ak
                                          08 09
1 2 3
               04 5 06
VAR G1=4-1 6-3/7/8-1 9-3/10
:del 7
:dis sia
                                                      Hy @10
\texttt{Cb} \backsim \texttt{G1} \backsim \texttt{Cb} \qquad \qquad \texttt{Ak} \backsim \texttt{Hy} \backsim \texttt{Ak}
                                         Hy∽ Ak
               04 5 06
                                          68 69
VAR G1=4-1 6-3/7/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
```

ECOUNT IS E4 C E1 O AT 5 ECOUNT IS E4 C E1 O AT 8 ECOUNT IS E6 C E2 O AT 10 29 ANSWERS

```
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
:VAR G1=4-1 6-3/8-1 9-3/10
G1 IN USE, CHANGE? (Y)/N:v
:dis sia
                                                    Hy @10
Cb G1 Cb Ak Hy Ak
1 2 3 04 5 06
                                       Hy∽ Ak
                                        08 09
VAR G1=4-1 6-3/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E4 C E1 O AT 5
ECOUNT IS E4 C E1 O AT 8
ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
:arr.dis
                                           Hy @10
Cb√G1√Cb
              Ak∽ Hy∽ Ak
                            Hy∽Ak
@8 @9
              04 5 06
1 2 3
VAR G1=4-1 6-3/8-1 9-3/10
· end
L11 STRUCTURE CREATED
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
               TRA L1 1- RN : 16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
             10 L3 AND >=2 46.150.18/RID
L4
             1 L3 AND OC4-C6-C6/ES
L5
             1 L3 AND C6-C6/ES
L6
               STR
L8
            29 L7
T.9
              SCR 1839
L10
            29 L7 AND L9
L11
               STR L7
```

=> >=2 46.150.18/rid and (oc4 or oc4-oc4)/es 16596122 RID.CNT >= 2 22121069 46.150.18/RID 11101089 >=2 46.150.18/RID (RID.CNT >= 2 (T) 46.150.18/RID) 1351114 OC4/ES 30009 OC4-OC4/ES

L12 336455 >=2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES

=> d sca

L12 336455 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN IN INDEX NAME NOT YET ASSIGNED MF C26 H28 F2 N2 O3

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1): HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L12 336455 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN IN INDEX NAME NOT YET ASSIGNED MF C23 H19 C13 N4 O5

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L12 336455 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN IN INDEX NAME NOT YET ASSIGNED MF C31 H33 F N6 08

Absolute stereochemistry. Double bond geometry as shown.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L12 336455 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN INDEX NAME NOT YET ASSIGNED IN

C23 H20 C12 N4 O5 MF

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

L2

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON L1 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

L3 16 SEA L2 10 L3 AND >=2 46.150.18/RID L4

L5 1 L3 AND OC4-C6-C6/ES L6 1 L3 AND C6-C6/ES

L7 STR L8 29 L7

L9 SCR 1839 29 L7 AND L9 L10

L11 STR L7

```
L12 336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
=> d que 111
               STR
                                         Hy @10
Cb/G1/Cb
           Ak∽Hy∽Ak
@4 5 @6
                           Hy∽Ak
@8 @9
1 2 3
VAR G1=4-1 6-3/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E4 C E1 O AT 5
ECOUNT IS E4 C E1 O AT 8
ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
    FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
              SET AUTOSEARCH ON
             1 US20050249857/PN
L1
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
              TRA L1 1- RN :
                                 16 TERMS
   FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
            16 SEA L2
L4
            10 L3 AND >=2 46.150.18/RID
L5
            1 L3 AND OC4-C6-C6/ES
L6
            1 L3 AND C6-C6/ES
L7
              STR
L8
            29 T.7
L9
              SCR 1839
L10
            29 L7 AND L9
L11
               STR I.7
L12 336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
=> 111 sub=112 sam
SAMPLE SUBSET SEARCH INITIATED 14:43:57 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED - 16809 TO ITERATE
11.9% PROCESSED 2000 ITERATIONS
                                                            50 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): 328415 TO 343945
PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET):
                                                    8112 TO
                                                               10714
```

50 SEA SUB=L12 SSS SAM L11

=> d sca

L13 50 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

Benzeneacetonitrile, 2-fluoro-a-[[5-(2-nitrophenyl)-2furanvllmethvlenel-

C19 H11 F N2 O3

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L13 50 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN IN Benzenepropanenitrile, 3-methyl-α-[[5-(3-nitrophenyl)-2-

furanyl]methylene]-β-oxo-

C21 H14 N2 O4 MF

$$\begin{array}{c|c} & NC & 0 \\ \hline \\ O_2N & CH = C-C \\ \end{array}$$

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON

1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

16 SEA L2 T. 4

10 L3 AND >=2 46.150.18/RID

L5 1 L3 AND OC4-C6-C6/ES

```
1 L3 AND C6-C6/ES
L6
1.7
                STR
L8
              29 L7
L9
                 SCR 1839
L10
              29 L7 AND L9
L11
                STR L7
L12
        336455 >=2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
              50 L11 SAM SUB=L12
=> str 111
:dis
                                              Hy @10
               Ak ~ Hy ~ Ak
@4 5 @6
                            Hy∽Ak
@8 @9
Chv/G1v/Ch
1 2 3
VAR G1=4-1 6-3/8-1 9-3/10
:eco 4 6 9 m1-x2 c,dis sia
                                              Hy @10
Cb G1 Cb
               Ak√ Hv√ Ak
                                 Hy∽Ak
1 2 3
               @4 5 @6
                                 68 69
VAR G1=4-1 6-3/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M1-X2 C AT 4
ECOUNT IS E4 C E1 O AT 5
ECOUNT IS M1-X2 C AT 6
ECOUNT IS E4 C E1 O AT 8
ECOUNT IS M1-X2 C AT 9
ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
:dis sia
                                              Hv @10
Cb\G1\Cb Ak\Hy\Ak
1 2 3 @4 5 @6
                             Hy∽Ak
@8 @9
VAR G1=4-1 6-3/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
DEFAULT ECLEVEL IS LIMITED ECOUNT IS MI-X2 C AT 4 ECOUNT IS E4 C E1 O AT 5 ECOUNT IS MI-X2 C AT 6 ECOUNT IS E4 C E1 O AT 8 ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
```

STEREO ATTRIBUTES: NONE

```
:end
L14 STRUCTURE CREATED
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
    FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
             1 US20050249857/PN
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
T.2
               TRA L1 1- RN : 16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
            16 SEA L2
L4
            10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
L6
L7
               STR
L8
            29 L7
1.9
               SCR 1839
T-10
            29 L7 AND L9
L11
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
            50 L11 SAM SUB=L12
L13
L14
               STR L11
=> 114 sub=112 sam
SAMPLE SUBSET SEARCH INITIATED 14:45:47 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED - 16809 TO ITERATE
11.9% PROCESSED
                   2000 ITERATIONS
                                                              32 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): 328415 TO 343945
PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET):
                                                     4395 TO
L15
           32 SEA SUB=L12 SSS SAM L14
=> d sca
L15 32 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
     2(3H)-Furanone, 4-[(3,4-dimethoxyphenyl)methyl]dihydro-3-hydroxy-3-[(4-
     hydroxy-3,5-dimethoxyphenyl)methyl]-, (3S,4S)-
ME
    C22 H26 O8
Absolute stereochemistry. Rotation (-).
```

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

- L15 32 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN 2(3H)-Furanone, dihydro-3-[[3-methoxy-4,5-bis(phenylmethoxy)phenyl]methyl]-4-[[3-methoxy-4-(phenylmethoxy)phenyl]methyl]-
- MF C41 H40 07

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

- L15 32 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN 2,4(1H,3H)-Pyrimidinedione, 1-[(2Z)-2-[5-oxo-3,4-bis(2-phenylacety1)-2(5H)-furanylidene]ethyl]-5-[2-(4-pentylphenyl)ethynyl]- C39 H34 N2 06

Double bond geometry as shown.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L15 32 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Furan, 3-fluoro-5-hexyl-2-phenyl-4-[(1E)-2-phenylethenyl]-MF C24 H25 F O

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

- L15 32 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN 2(3H)-Furanone, 4-[(3,4-dimethoxyphenyl)hydroxymethyl]dihydro-5-methoxy-3-[(2,4,5-trimethoxyphenyl)methylene]-
- MF C24 H28 O9

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON

L1 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008

L2 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

L3 16 SEA L2

L4 10 L3 AND >=2 46.150.18/RID

L5 1 L3 AND OC4-C6-C6/ES

L6 1 L3 AND C6-C6/ES

L7 STR L8 29 L7

L9 SCR 1839

L10 29 L7 AND L9

L11 STR L7 L12 336455 >=2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES

L13 50 L11 SAM SUB=L12 L14 STR L11

L15 32 L14 SAM SUB=L12

=> e furan/cn

E1 1 FURAMPICILLIN SODIUM/CN

E2 1 FURAMTERENE/CN E3 1 --> FURAN/CN

E4 1 FURAN 1/CN

E5 1 FURAN 2/CN

E6 1 FURAN 2 (DYE)/CN
E7 1 FURAN 2 (FISH MEDICATION)/CN

E8 1 FURAN 6/CN

E9 1 FURAN CATION/CN

E10 1 FURAN CATION RADICAL/CN

E11 1 FURAN CONJUGATE MONOACID/CN E12 1 FURAN ENDO-PEROXIDE/CN

=> e3

L16 1 FURAN/CN

=> d str

L16 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2008 ACS on STN



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

| => e thf/cn | | |
|-------------|----|----------------------------|
| => e thi/ch | | |
| E1 | 1 | THEXYLTRICHLOROSILANE/CN |
| E2 | 1 | THEXYLTRIMETHOXYSILANE/CN |
| E3 | 1> | THF/CN |
| E4 | 1 | THF 10/CN |
| E5 | 1 | THF 451-10A/CN |
| E6 | 1 | THF CATION RADICAL/CN |
| E7 | 1 | THF CYCLIC DIMER/CN |
| E8 | 1 | THF CYCLIC OCTAMER/CN |
| E9 | 1 | THF CYCLIC TETRAMER/CN |
| E10 | 1 | THF HEPTADECAHYDRATE/CN |
| E11 | 1 | THF HOMOPOLYMER/CN |
| E12 | 1 | THF HOMOPOLYMER ACETATE/CN |

=> e3

=> d str rsd

L17 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2008 ACS on STN



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

Ring System Data

| Elementa | l Elementa | 1 Size | of Ring | System Rir | ig RID |
|----------|------------|---------|-----------|--------------|-----------------|
| Analysis | Sequence | the F | ings For | mula Identi | fier Occurrence |
| EA | l ES | SZ | R | F RII | Count |
| | -+ | -+ | + | + | + |
| C40 | IOC4 | 15 | IC40 | 16.138 | .1 1 |

```
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
L1
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
                TRA L1 1- RN :
                                   16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
L6
L7
                STR
L8
             29 L7
L9
                SCR 1839
T-10
             29 L7 AND L9
L11
               STR L7
L12
         336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
             50 L11 SAM SUB=L12
L13
L14
                STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
             1 E3
              E THF/CN
L17
             1 E3
=> 112 and 16.138.1
            0 16.138.1
L18
             0 L12 AND 16.138.1
=> del 118 y
=> 112 and 16.138.1/rid
        554802 16.138.1/RID
1.18
       130953 L12 AND 16.138.1/RID
=> d sca
L18 130953 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
     INDEX NAME NOT YET ASSIGNED
IN
MF
     C43 H50 N3 O7 P
```

Absolute stereochemistry.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

```
(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
```

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON

L1 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 L2 TRA L1 1- RN : 16 TERMS

```
FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
```

L3 16 SEA L2

L4 10 L3 AND >=2 46.150.18/RID

L5 1 L3 AND OC4-C6-C6/ES

L6 1 L3 AND C6-C6/ES

L7 STR

L8 29 L7

L9 SCR 1839

L10 29 L7 AND L9 L11 STR L7

L12 336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES

L13 50 L11 SAM SUB=L12 L14 STR L11

L15 32 L14 SAM SUB=L12

E FURAN/CN

L16 1 E3

E THF/CN

L17 1 E3

L18 130953 L12 AND 16.138.1/RID

=> 114 sub=118 sam

SAMPLE SUBSET SEARCH INITIATED 14:48:08 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED - 6522 TO ITERATE

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): 125598 TO 13592
PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET): 2437 TO 3953

L19 49 SEA SUB=L18 SSS SAM L14

=> d sca

L19 49 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Methanone, 1,1'-[(3R,4R)-tetrahydro-3,4-furandiy1]bis[1-(3,4-dimethoxypheny1)-

MF C22 H24 07

Absolute stereochemistry. Rotation (+).

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L19 49 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN 2(3H)-Furanone, dihydro-3-pentyl-5-phenyl-4-(phenylmethyl)-, (3R,4R,5S)-rel-

MF C22 H26 O2

Relative stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1): HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

```
(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
                SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
                TRA L1 1- RN :
                                     16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
1.3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
              1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
L6
L7
               STR
L8
             29 L7
                SCR 1839
L9
L10
             29 L7 AND L9
L11
                STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB=L12
L14
                STR L11
L15
             32 L14 SAM SUB=L12
                E FURAN/CN
L16
              1 E3
               E THF/CN
L17
             1 E3
L18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
=> str 114
:dis
              Ak∽ Hy∽ Ak
                                            Hy @10
                           Hy^Ak
08 09
Cb G1 Cb
               04 5 06
1 2 3
VAR G1=4-1 6-3/8-1 9-3/10
:att o 1 se.3 o se.dis sia
                            Ak√ Hy√ Ak
               Hy√ Ak
                                     11 1 2 3 12
@4 5 @6
                08 09
VAR G1=4-1 6-3/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M1-X2 C AT 4
ECOUNT IS E4 C E1 O AT 5
ECOUNT IS E4 C E1 O AT 6
ECOUNT IS E4 C E1 O AT 8
ECOUNT IS M1-X2 C AT 9
ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 11
```

STEREO ATTRIBUTES: NONE

:FIL STNGUIDE

DATA TYPED NOT VALID

You entered an unknown command name or an incomplete command. For more information, enter "HELP" and the command name. Enter "HELP" to see a list of all the commands. :end

L20 STRUCTURE CREATED

=> b stng

COST IN U.S. DOLLARS

SINCE FILE

TOTAL ENTRY SESSION 94.40 112.31

> TOTAL SESSION

112.67

FULL ESTIMATED COST

FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008 USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

FILE CONTAINS CURRENT INFORMATION. LAST RELOADED: Jun 6, 2008 (20080606/UP).

=> b rea

COST IN U.S. DOLLARS

SINCE FILE ENTRY

0.36

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.

PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 11 JUN 2008 HIGHEST RN 1027436-61-8 DICTIONARY FILE UPDATES: 11 JUN 2008 HIGHEST RN 1027436-61-8

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 9, 2008.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

```
FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
               TRA L1 1- RN : 16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
             10 L3 AND >=2 46.150.18/RID
L4
L5
             1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
L6
L7
               STR
L8
            29 L7
L9
               SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
L12
       336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
            50 L11 SAM SUB=L12
L13
L14
               STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
L17
              1 E3
L18
       130953 L12 AND 16.138.1/RID
T.19
            49 L14 SAM SUB=L18
T-20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
=> str 120
:dis
                          Hy @10
              Hv∽ Ak
                                    0-Cb-G1-Cb-0
Ak√ Hv√ Ak
@4 5 @6
              e8 e9
                                    11 1 2 3 12
VAR G1=4-1 6-3/8-1 9-3/10
:eco 1 3 e6 c,qqc 1 3 mcv,dis sia
                           Hv @10
Ak∽ Hy∽ Ak
               Hv√ Ak
                                     0-Cb\sigma G1\sigma Cb-0
04 5 06
               08 09
                                     11 1 2 3 12
VAR G1=4-1 6-3/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS MCY AT 1
GGCAT IS MCY AT 3
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E6 C AT 1
ECOUNT IS E6 C AT 3
ECOUNT IS M1-X2 C AT 4
ECOUNT IS E4 C E1 O AT
ECOUNT IS M1-X2 C AT 6
ECOUNT IS E4 C E1 O AT 8
ECOUNT IS M1-X2 C AT 9
ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
```

RING(S) ARE ISOLATED OR EMBEDDED

```
NUMBER OF NODES IS 11
STEREO ATTRIBUTES: NONE
:end
L21 STRUCTURE CREATED
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
                TRA L1 1- RN :
                                    16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
              1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
1.6
1.7
               STR
L8
             29 L7
L9
               SCR 1839
L10
             29 L7 AND L9
               STR L7
L11
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
            50 L11 SAM SUB=L12
L14
               STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
L17
              1 E3
L18
         130953 L12 AND 16.138.1/RID
L19
             49 L14 SAM SUB=L18
L20
                STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
                STR 1.20
=> scr 1707 or 1708
L22 SCREEN CREATED
=> scr 1840
L23 SCREEN CREATED
=> dis sia
L23 HAS NO ANSWERS
L23
               SCR 1840
=> d his
```

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

```
FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
             1 US20050249857/PN
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
               TRA L1 1- RN :
                                   16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
T.3
            16 SEA L2
L4
            10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
L6
L7
               STR
            29 L7
L8
L9
               SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
L12
       336455 >=2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
            50 L11 SAM SUB=L12
L13
L14
               STR L11
L15
            32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
             1 E3
L18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
=> 122 and 123 and 121
SAMPLE SEARCH INITIATED 15:00:39 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 431505 TO ITERATE
 0.5% PROCESSED
                   2000 ITERATIONS
                                                                2 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**
                        BATCH **INCOMPLETE**
PROJECTED ITERATIONS:
                         8592492 TO 8667708
PROJECTED ANSWERS:
                             7384 TO 9876
             2 SEA SSS SAM L22 AND L23 AND L21
L24
=> d sc
'SC' IS NOT A VALID FORMAT FOR FILE 'REGISTRY'
The following are valid formats:
```

Substance information can be displayed by requesting individual

fields or predefined formats. The predefined substance formats are: (RN = CAS Registry Number) REG RN - Index Name, MF, and structure - no RN SAM FIDE - All substance data, except sequence data - FIDE, but only 50 names SOIDE - IDE, plus sequence data SOIDE3 - Same as SOIDE, but 3-letter amino acid codes are used SOD - Protein sequence data, includes RN SQD3 - Same as SQD, but 3-letter amino acid codes are used SON - Protein sequence name information, includes RN CALC - Table of calculated properties EPROP - Table of experimental properties PROP - EPROP and CALC Any CA File format may be combined with any substance format to obtain CA references citing the substance. The substance formats must be cited first. The CA File predefined formats are: ABS -- Abstract APPS -- Application and Priority Information BIB -- CA Accession Number, plus Bibliographic Data CAN -- CA Accession Number CBIB -- CA Accession Number, plus Bibliographic Data (compressed) IND -- Index Data IPC -- International Patent Classification PATS -- PI, SO STD -- BIB, IPC, and NCL IABS -- ABS, indented, with text labels IBIB -- BIB, indented, with text labels ISTD -- STD format, indented OBIB ----- AN, plus Bibliographic Data (original) OIBIB ----- OBIB, indented with text labels SBIB ----- BIB, no citations SIBIB ----- IBIB, no citations The ALL format gives FIDE BIB ABS IND RE, plus sequence data when it is available. The MAX format is the same as ALL. The IALL format is the same as ALL with BIB ABS and IND indented, with text labels. For additional information, please consult the following help messages:

HELP DFIELDS -- To see a complete list of individual display fields. HELP FORMATS -- To see detailed descriptions of the predefined formats. ENTER DISPLAY FORMAT (IDE):end

=> d sca

L24 2 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN IN 3(2H)-Furanone, 4-[(3,4-dimethoxyphenyl)methyl]-2-(4-hydroxy-3methoxyphenyl)-

ME C20 H20 O6

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

L1

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008

SET AUTOSEARCH ON

1 US20050249857/PN FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

TRA L1 1- RN : 16 TERMS

L2

L3 16 SEA L2

L4 10 L3 AND >=2 46.150.18/RID

L5 1 L3 AND OC4-C6-C6/ES

L6 1 L3 AND C6-C6/ES

L7 STR

L8 29 L7

L9 SCR 1839

L10 29 L7 AND L9 Lli STR L7

336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES L12 L13 50 L11 SAM SUB=L12

L14 STR L11

32 L14 SAM SUB=L12 L15

E FURAN/CN L16 1 E3

E THF/CN

1 E3 1.18 130953 L12 AND 16.138.1/RID

L19 49 L14 SAM SUB=L18

L20 STR L14 FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008

FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008

L21 STR L20

L22 SCR 1707 OR 1708 L23 SCR 1840

L23 SCR 1840 L24 2 L22 AND L23 AND L21

=> 122 and 123 and 121 sub=118 sam

SAMPLE SUBSET SEARCH INITIATED 15:01:14 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED - 4728 TO ITERATE

42.3% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET):

PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): 90437 TO 98683

36 ANSWERS

2255

1149 TO

L25 36 SEA SUB=L18 SSS SAM L22 AND L23 AND L21

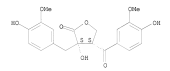
=> d sca

L25 36 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

N 2(3H)-Furanone, dihydro-3-hydroxy-4-(4-hydroxy-3-methoxybenzoyl)-3-[(4-hydroxy-3-methoxyphenyl)methyl]-, (3S, 4S)-

MF C20 H20 O8

Absolute stereochemistry. Rotation (+).



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

- L25 36 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN Benzenepropanoic acid, 2-(phenylmethoxy)-4-[[(3R,4R)-tetrahydro-2-oxo-4-[[3-(phenylmethoxy)phenyl]methyl]-3-furanyl]methyl]-, rel-

MF C35 H34 O6

Relative stereochemistry.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

- L25 36 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN 2(3H)-Furanone, 4-[(R)-(3,4-dihydroxyphenyl)hydroxymethyl]dihydro-3-[(4-hydroxy-3-methoxyphenyl)methyl]-, (3R,4R)-

MF C19 H20 07

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

- L25 36 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN 2(3H)-Furanone, dihydro-3-[[3-methoxy-4,5-bis(phenylmethoxy)phenyl]methyl]-
- 4-[[3-methoxy-4-(phenylmethoxy)phenyl]methyl]-MF C41 H40 O7

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L25 36 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
10 2(3H)-Furanone, dihydro-4-(hydroxymethyl)-5-[3-methoxy-4-(phenylmethoxy)phenyl]-3-[[3-methoxy-4-(phenylmethoxy)phenyl]methyl]-,
135, 4R, 5R)MF C34 H34 07

Absolute stereochemistry. Rotation (-).

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

```
FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
               TRA L1 1- RN : 16 TERMS
L2
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
            16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
L7
               STR
L8
            29 L7
L9
               SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
L12
       336455 >=2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
            50 L11 SAM SUB=L12
L14
               STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
              1 E3
L18
        130953 L12 AND 16.138.1/RID
T.19
            49 L14 SAM SUB=L18
T-20
               STR 1.14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
             2 L22 AND L23 AND L21
L25
            36 L22 AND L23 AND L21 SAM SUB=L18
=> str 121
:dis
                          Hy 010 O-Cb~G1~Cb-O
              Hv∽ Ak
Ak√ Hv√ Ak
@4 5 @6
               e8 e9
                                     11 1 2 3 12
VAR G1=4-1 6-3/8-1 9-3/10
:att c1 11,c1 12,dis sia
                           Hy @10 C~~O—Cb~G1~Cb—O~~C
Ak∽ Hy∽ Ak
              Hy∽ Ak
@4 5 @6
               ଜନି ଜନ
                                    13 11 1 2 3 12 14
VAR G1=4-1 6-3/8-1 9-3/10
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
GGCAT IS MCY AT 1
GGCAT IS MCY AT 3
GGCAT IS PCY AT 10
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E6 C AT 1
ECOUNT IS E6 C AT 3
ECOUNT IS M1-X2 C AT 4
ECOUNT IS E4 C E1 O AT
ECOUNT IS M1-X2 C AT 6
ECOUNT IS E4 C E1 O AT 8
```

ECOUNT IS M1-X2 C AT 9 ECOUNT IS E6 C E2 O AT 10

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 13

STEREO ATTRIBUTES: NONE

:att o 13 vn d.att o 14 vn d.dis sia

16 15 Hy @10 Hv√Ak Ak ~ Hv ~ Ak @4 5 @6 68 R9 13 11 1 2 3 12

VAR G1=4-1 6-3/8-1 9-3/10 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM GGCAT IS MCY AT 1 IS MCY AT GGCAT IS PCY AT 10 GGCAT DEFAULT ECLEVEL IS LIMITED ECOUNT IS E6 C AT 1 ECOUNT IS E6 C AT 3 ECOUNT IS M1-X2 C AT 4 ECOUNT IS E4 C E1 O AT ECOUNT IS M1-X2 C AT ECOUNT IS E4 C E1 O AT 8 ECOUNT IS M1-X2 C AT 9 ECOUNT IS E6 C E2 O AT 10

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 15

STEREO ATTRIBUTES: NONE

:d his

DATA TYPED NOT VALID

You entered an unknown command name or an incomplete command. For more information, enter "HELP" and the command name. Enter "HELP" to see a list of all the commands.

:dis sia

16 Hv @10 Ak∽ Hy∽ Ak Hv∽ Ak @4 5 @6 ดลี ดจ ~~0—Cb√G1√Cb—0~√C 13 11 1 2 3 12

VAR G1=4-1 6-3/8-1 9-3/10 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM GGCAT IS MCY AT 1 GGCAT IS MCY AT 3 GGCAT IS PCY AT 10 DEFAULT ECLEVEL IS LIMITED ECOUNT IS E6 C AT 1 ECOUNT IS E6 C AT 3 ECOUNT IS M1-X2 C AT 4

```
ECOUNT IS E4 C E1 O AT 5
ECOUNT IS M1-X2 C AT 6
ECOUNT IS E4 C E1 O AT 8
ECOUNT IS M1-X2 C AT 9
ECOUNT IS E6 C E2 O AT 10
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 15
STEREO ATTRIBUTES: NONE
:end
L26 STRUCTURE CREATED
=> d hjis
L26 HAS NO ANSWERS
'HJIS ' IS NOT A VALID STRUCTURE FORMAT KEYWORD
Structure Formats
SIA ---- Structure Image, Attributes, and map table if it contains
         data. (Default)
SIM ---- Structure IMage.
SAT ---- Structure ATtributes and map table if it contains data.
SCT ---- Structure Connection Table and map table if it contains
         data.
SDA ---- All Structure DAta (image, attributes, connection table and
         map table if it contains data).
NOS ---- NO Structure data.
ENTER STRUCTURE FORMAT (SIA), SCT, SDA, SIM, SAT, NOS:end
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
    FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
               TRA L1 1- RN :
                                    16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
1.3
            16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
L7
               STR
             29 L7
L8
               SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
            50 L11 SAM SUB=L12
L14
               STR L11
L15
            32 L14 SAM SUB=L12
              E FURAN/CN
1.16
             1 E3
              E THF/CN
             1 E3
```

```
1.18
        130953 L12 AND 16.138.1/RID
1.19
             49 L14 SAM SUB=L18
L20
               STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
=> 126 and 122 and 123 sub=118 sam
SAMPLE SUBSET SEARCH INITIATED 15:03:09 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED - 1302 TO ITERATE
100.0% PROCESSED
                   1302 ITERATIONS
                                                                 4 ANSWERS
SEARCH TIME: 00.00.01
PROJECTIONS (WITHIN SPECIFIED SUBSET):
                                                ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET):
                                                      23876 TO
                                                                28204
PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET):
                                                           4 TO
             4 SEA SUB=L18 SSS SAM L26 AND L22 AND L23
L27
=> d sca
```

L27 4 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN 2(3H)-Furanone, 4-[(acetyloxy)[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-TN [(acetyloxy)[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-ME C28 H28 O12

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

4 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN TN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,Z)- (9CI) C24 H20 O9 ME

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

- L27 4 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
- IN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,E)- (9CI)
- MF C24 H20 O9

Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> 126 and 122 and 123 sub=118 full FULL SUBSET SEARCH INITIATED 15:03:33 FILE 'REGISTRY' FULL SUBSET SCREEN SEARCH COMPLETED - 25929 TO ITERATE

100.0% PROCESSED 25929 ITERATIONS SEARCH TIME: 00.00.01 94 ANSWERS

SEARCH TIME: UU.UU.UI

L28

94 SEA SUB=L18 SSS FUL L26 AND L22 AND L23

:dis

```
=> sav tem q761c1/a 128
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
L1
              1 US20050249857/PN
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
               TRA L1 1- RN :
                                 16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
L7
               STR
L8
            29 L7
L9
               SCR 1839
T-10
            29 L7 AND L9
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
            50 L11 SAM SUB=L12
L14
               STR L11
L15
            32 L14 SAM SUB=L12
               E FURAN/CN
             1 E3
L16
               E THF/CN
L17
             1 E3
L18
        130953 L12 AND 16.138.1/RID
L19
             49 L14 SAM SUB=L18
L20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
               SCR 1840
L23
L24
             2 L22 AND L23 AND L21
L25
            36 L22 AND L23 AND L21 SAM SUB=L18
L26
               STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
            94 L26 AND L22 AND L23 FULL SUB=L18
               SAV TEM G761C1/A L28
=> str 17
:dis
                              Ak @7
                                                   Hy @10
Cbv^G1v^Cb
             Ak√ Hy√ Ak
                                      Hv√ Ak
              04 5 06
                                        08 09
VAR G1=4-1 6-3/7/8-1 9-3/10
:del s 4,s 8,10
```

```
Ak @7
Cb√G1√Cb
VAR G1=4-1 6-3/7/8-1 9-3/10
:del 7
:nod 2 ak.eco 2 m3-x5 c.dis sia
Cb\sigma Ak\sigma Cb
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M3-X5 C AT 2
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 3
STEREO ATTRIBUTES: NONE
:att c2 1,c2 3
:dis sia
C^-C^-Cb^Ak^Cb-^C-^C
5 4 1 2 3 6 7
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M3-X5 C AT 2
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 7
STEREO ATTRIBUTES: NONE
:nod 5 7 o,att o 4 vn d,6 o vn d,dis sia
   0
0~~C~~Cb~Ak~Cb~~C~~O
5 4 1 2 3 6
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M3-X5 C AT 2
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
:del 8 9
:dis
```

```
0~-C~-Cb~ Ak~ Cb~~ C~~ 0
5 4 1 2 3 6 7
:nod 4 6 0,5 7 c,att o 5 vn d,7 o vn d,dis sia
                        9
 0
                        0
5 C~~O~~Cb~ Ak~ Cb~~O~~ C
       1 2 3
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS M3-X5 C AT 2
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
:end
L29 STRUCTURE CREATED
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
                TRA L1 1- RN :
                                    16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
L7
               STR
L8
             29 L7
               SCR 1839
L9
L10
             29 L7 AND L9
L11
                STR L7
L12
         336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB-L12
L14
                STR L11
             32 L14 SAM SUB=L12
L15
                E FURAN/CN
L16
              1 E3
              E THF/CN
              1 E3
1.18
         130953 L12 AND 16.138.1/RID
            49 L14 SAM SUB=L18
L19
L20
                STR L14
```

```
FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
```

FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
STR L20
L22 SCR 1707 OR 1708
L23 SCR 1840
L24 2 L22 AND L23 AND L21
L25 36 L22 AND L23 AND L21 SAM SUB=L18

L25 36 L22 AND L23 AND L21 SAM SUB=L18 L26 STR L21 L27 4 L26 AND L22 AND L23 SAM SUB=L18

L28 94 L26 AND L22 AND L23 FULL SUB=L18 SAV TEM G761C1/A L28 L29 STR L7

=> 129

=> 129
SAMPLE SEARCH INITIATED 15:06:34 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 159857 TO ITERATE

SAMPLE SCREEN SEARCH COMPLETED - 159857 TO ITERATIONS

1.3% PROCESSED 2000 ITERATIONS

INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**
PROJECTED ITERATIONS: 3173552 TO 3220728
PROJECTED ANSWERS: 2439 TO 3955

L30 2 SEA SSS SAM L29

=> d sca

L30 2 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN 1,4-Benzenedicarboxylic acid, polymer with 4-(acetyloxy)benzoic acid, bis[2-(acetyloxy)ethyl] 1,4-benzenedicarboxylate, 1,2-ethanediol and (1-methylethylidene)di-4,1-phenylene diacetate, block (9C1)

2 ANSWERS

MF (C19 H20 O4 . C16 H18 O8 . C9 H8 O4 . C8 H6 O4 . C2 H6 O2)x

CI PMS

CM 1

CM 2

L7 STR L8 29 L7 L9 SCR 1839 29 L7 AND L9 L10

L410 L3 AND >=2 46.150.18/RID 1 L3 AND OC4-C6-C6/ES L6 1 L3 AND C6-C6/ES

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 L3 16 SEA L2

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

1 US20050249857/PN

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

=> d his

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

HO₂C

L1

L2

5

HO-CH2-CH2-OH CM

CM

CM 3

CO₂H

```
STR L7
1.12
         336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB=L12
L14
               STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
L17
             1 E3
L18
         130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
                STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
                STR L20
L22
                SCR 1707 OR 1708
L23
                SCR 1840
L24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
L27
              4 L26 AND L22 AND L23 SAM SUB=L18
L28
             94 L26 AND L22 AND L23 FULL SUB=L18
                SAV TEM G761C1/A L28
L29
                STR L7
L30
             2 L29
=> str 129
:dis
 8
                        9
5 C~-O~-Cb~Ak~ Cb~~O~~C
           2 3
:nod 2 q1
:rep g1=(3-5) c,dis sia
 8
 0
5 C~-0~-Cb~G1~Cb~~0~~C
           2 3
REP G1=(3-5) C
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
:end
L31 STRUCTURE CREATED
=> d his
```

```
(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
    FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
             1 US20050249857/PN
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
               TRA L1 1- RN :
                                   16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
1.3
            16 SEA L2
L4
            10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
L7
               STR
L8
            29 L7
L9
               SCR 1839
            29 L7 AND L9
L10
L11
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
            50 L11 SAM SUB=L12
T-14
               STR L11
L15
            32 L14 SAM SUB=L12
               E FURAN/CN
L16
             1 E3
              E THF/CN
             1 E3
L17
L18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
               SCR 1707 OR 1708
L22
L23
               SCR 1840
L24
             2 L22 AND L23 AND L21
L25
            36 L22 AND L23 AND L21 SAM SUB=L18
L26
               STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
            94 L26 AND L22 AND L23 FULL SUB=L18
               SAV TEM G761C1/A L28
L29
               STR L7
L30
             2 L29
L31
               STR L29
=> 131
SAMPLE SEARCH INITIATED 15:07:17 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 132971 TO ITERATE
 1.5% PROCESSED
                   2000 ITERATIONS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
```

FULL FILE PROJECTIONS: ONLINE **INCOMPLETE**
BATCH **INCOMPLETE**

SEARCH TIME: 00.00.01

0 ANSWERS

PROJECTED ITERATIONS: 2637845 TO 2680995 PROJECTED ANSWERS: 0 TO

L32 0 SEA SSS SAM L31

=> str 131 :dis

REP G1=(3-5) C :nod 2 ak.dis sia

ELEMENT SYMBOL NOT VALID

An element symbol specified is not valid. Enter "HELP NODE" for more information. :dis sia

8 9 ~~ 0 ~~ Cb ~ G1 ~ Cb ~~ 0 ~~ C , 4 1 2 3 6

REP G1=(3-5) C NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 9

STEREO ATTRIBUTES: NONE :nod 2 ak, dis sia

$$\begin{picture}(60,0)(0,0) \put(0,0){\line(0,0){10}} \put(0,0){\line(0,0){10}}$$

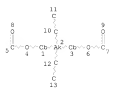
NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 9

STEREO ATTRIBUTES: NONE :att c2 2 vn,c2 2 vs,dis



:nod 11 13 o,dis sia

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 13

STEREO ATTRIBUTES: NONE :end L33 STRUCTURE CREATED

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON L1 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 3 16 SEA L2

L6 1 L3 AND C6-C6/ES

```
STR
1.8
             29 L7
L9
                SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
         336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB=L12
L14
                STR L11
L15
             32 L14 SAM SUB=L12
                E FURAN/CN
L16
              1 E3
               E THF/CN
              1 E3
L18
         130953 L12 AND 16.138.1/RID
L19
             49 L14 SAM SUB=L18
                STR L14
L20
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
                STR L20
L21
                SCR 1707 OR 1708
L22
L23
                SCR 1840
L24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
L27
              4 L26 AND L22 AND L23 SAM SUB=L18
             94 L26 AND L22 AND L23 FULL SUB=L18
L28
                SAV TEM G761C1/A L28
L29
                STR L7
L30
              2 L29
L31
                STR L29
L32
              0 L31
L33
                STR L31
=> 133
GENERIC GROUP NOT VALID HERE
Generic groups may not be used in these circumstances:
       1. Any generic group node (e.g., Hy) in a ring.
       An Ak node attached to another Ak node.
       3. An Ak node with three or more attachments where
           one or more of the attachments is to a C node.
=> str 133
:dis
           11
            0
 8
         10 c
        Cb~Ak~Cb~~O
         1
                3
         12 C
```

13

```
:end
L34 STRUCTURE CREATED
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
L1
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
T. 2
               TRA L1 1- RN : 16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
             1 L3 AND C6-C6/ES
L6
L7
               STR
L8
             29 L7
L9
               SCR 1839
T-10
             29 L7 AND L9
L11
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
            50 L11 SAM SUB=L12
L14
               STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
L17
             1 E3
L18
         130953 L12 AND 16.138.1/RID
L19
             49 L14 SAM SUB=L18
L20
                STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
                STR L20
L22
                SCR 1707 OR 1708
L23
               SCR 1840
1.24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
             94 L26 AND L22 AND L23 FULL SUB=L18
                SAV TEM G761C1/A L28
L29
               STR L7
L30
             2 L29
L31
               STR L29
L32
             0 L31
               STR L31
L34
               STR L33
=> d que 133
L33
               STR
```

L18

```
11
            0
 8
                         9
          10 c
 0
         Cb Ak
          12 C
           13
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 13
STEREO ATTRIBUTES: NONE
=> del 133- v
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
                SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
                TRA L1 1- RN :
                                     16 TERMS
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
              1 L3 AND OC4-C6-C6/ES
              1 L3 AND C6-C6/ES
L6
L7
                STR
L8
             29 L7
L9
                SCR 1839
             29 L7 AND L9
L11
                STR L7
         336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L12
L13
             50 L11 SAM SUB=L12
L14
                STR L11
             32 L14 SAM SUB=L12
                E FURAN/CN
L16
              1 E3
                E THF/CN
              1 E3
```

130953 L12 AND 16.138.1/RID

```
1.19
           49 L14 SAM SUB=L18
L20
               STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
             94 L26 AND L22 AND L23 FULL SUB=L18
                SAV TEM G761C1/A L28
L29
                STR L7
L30
             2 L29
L31
               STR L29
L32
             0 L31
=> d aue 131
L31
                STR
 8
5 C~-0~ Cb~ G1~ Cb~~ 0~~ C
REP G1=(3-5) C
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS
STEREO ATTRIBUTES: NONE
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
                SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
    FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
                TRA L1 1- RN : 16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
T.3
             16 SEA L2
T. 4
             10 L3 AND >=2 46.150.18/RID
1.5
             1 L3 AND OC4-C6-C6/ES
1.6
             1 L3 AND C6-C6/ES
L7
               STR
```

```
1.8
            29 L7
1.9
               SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB=L12
L14
               STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
              1 E3
L18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
                STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
1.21
                STR L20
L22
                SCR 1707 OR 1708
L23
                SCR 1840
L24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
1.26
                STR L21
             4 L26 AND L22 AND L23 SAM SUB=L18
L27
L28
             94 L26 AND L22 AND L23 FULL SUB=L18
                SAV TEM G761C1/A L28
L29
               STR L7
L30
             2 L29
L31
               STR L29
L32
             0 L31
=> scr 1701 or 1702 or 1703 or 1704 or 1705
L33 SCREEN CREATED
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
                SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
                TRA L1 1- RN : 16 TERMS
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
             10 L3 AND >=2 46.150.18/RID
L4
L5
             1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
               STR
1.8
             29 L7
T.9
               SCR 1839
T.10
             29 L7 AND L9
               STR L7
L12
       336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
```

```
50 L11 SAM SUB=L12
1.13
L14
              STR L11
L15
            32 L14 SAM SUB=L12
              E FURAN/CN
L16
             1 E3
              E THF/CN
L17
             1 E3
L18
       130953 L12 AND 16.138.1/RID
L19
           49 L14 SAM SUB=L18
L20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
T.21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
             2 L22 AND L23 AND L21
L25
            36 L22 AND L23 AND L21 SAM SUB=L18
L26
               STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
            94 L26 AND L22 AND L23 FULL SUB=L18
               SAV TEM G761C1/A L28
1.29
               STR L7
T.30
            2 L29
L31
              STR L29
L32
            0 L31
L33
               SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
=> 131 and 133
SAMPLE SEARCH INITIATED 15:12:17 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 29811 TO ITERATE
 6.7% PROCESSED
                  2000 ITERATIONS
                                                              0 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
FULL FILE PROJECTIONS: ONLINE **COMPLETE**
                       BATCH **COMPLETE**
                         585893 TO 606547
PROJECTED ITERATIONS:
PROJECTED ANSWERS:
                               0 TO
L34
             0 SEA SSS SAM L31 AND L33
=> 131 and 133 full
FULL SEARCH INITIATED 15:12:46 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 587831 TO ITERATE
98.8% PROCESSED 580727 ITERATIONS
                                                              33 ANSWERS
100.0% PROCESSED 587831 ITERATIONS
                                                              33 ANSWERS
SEARCH TIME: 00.00.25
L35
            33 SEA SSS FUL L31 AND L33
=> sav tem g761c1n/a 135
=> d sca
```

L35 33 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Propanoic acid, 2,2-dimethyl-, 3-(β-D-glucopyranosyloxy)-2-[3-[4-[[(2-methylpropoxy)carbonyl]oxy]phenyl]-1-oxopropyl]phenyl ester

MF C31 H40 O12

Absolute stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L35 33 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

10 4-Cyclohexene-1,3-dione, 6-(acetyloxy)-2-[(1635)-3-(acetyloxy)-5-[(2E)-3-[4-(acetyloxy)phenyl]-1-oxo-2-propenyl]-2,4-dihydroxy-6-oxo-3[(2R,38,4R,58,68)-3,4,5-tris(acetyloxy)-6-[(acetyloxy)methyl]tetrahydro-2Hpyran-2-yl]-1,4-cyclohexadien-1-yl]methylene]-4-[(2E)-3-[4(acetyloxy)phenyl]-1-oxo-2-propenyl]-5-hydrox-6-[(2S,3R,48,5R,68)-3,4,5tris(acetyloxy)-6-[(acetyloxy)methyl]tetrahydro-2H-pyran-2-yl]-, (2Z,6S)(9C1)

MF C67 H66 O34

Absolute stereochemistry. Double bond geometry as shown.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

1 US20050249857/PN

=> d his

L2

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 16 SEA L2

L3 16 SEA L2 L4 10 L3 AND >=2 46.150.18/RID

L5 1 L3 AND OC4-C6-C6/ES L6 1 L3 AND C6-C6/ES

L7 STR L8 29 L7

L9 SCR 1839 L10 29 L7 AND L9 L11 STR L7

L12 $336455 \ge 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES$

L13 50 L11 SAM SUB=L12 L14 STR L11

L15 32 L14 SAM SUB=L12

E FURAN/CN

L16 1 E3

```
E THE/CN
              1 E3
L18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
               STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
                STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
              2 L22 AND L23 AND L21
1.25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
             4 L26 AND L22 AND L23 SAM SUB=L18
L27
L28
            94 L26 AND L22 AND L23 FULL SUB=L18
               SAV TEM G761C1/A L28
L29
                STR L7
L30
             2 L29
L31
                STR L29
L32
             0 L31
L33
               SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L34
             0 L31 AND L33
L35
             33 L31 AND L33 FULL
                SAV TEM G761C1N/A L35
=> 128,135 and 13
L36
          0 (L28 OR L35) AND L3
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
L2
               TRA L1 1- RN :
                                   16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
1.3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
L6
              1 L3 AND C6-C6/ES
L7
               STR
             29 L7
L8
               SCR 1839
L9
             29 L7 AND L9
L10
L11
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
             50 L11 SAM SUB=L12
1.14
               STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
1.16
              1 E3
               E THF/CN
```

```
1 E3
1.18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
             2 L22 AND L23 AND L21
L25
            36 L22 AND L23 AND L21 SAM SUB=L18
L26
               STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
            94 L26 AND L22 AND L23 FULL SUB=L18
L28
               SAV TEM G761C1/A L28
L29
               STR L7
L30
             2 L29
L31
               STR L29
L32
             0 L31
L33
               SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L34
             0 L31 AND L33
L35
            33 L31 AND L33 FULL
               SAV TEM G761C1N/A L35
L36
             0 L28, L35 AND L3
=> b stng
                                                SINCE FILE
                                                                TOTAL
COST IN U.S. DOLLARS
                                                     ENTRY SESSION
FULL ESTIMATED COST
                                                    367.76
                                                              480.43
FILE 'STNGUIDE' ENTERED AT 15:14:39 ON 12 JUN 2008
USE IS SUBJECT TO THE TERMS OF YOUR CUSTOMER AGREEMENT
COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)
FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Jun 6, 2008 (20080606/UP).
=> b rea
COST IN U.S. DOLLARS
                                                SINCE FILE
                                                                TOTAL
                                                     ENTRY
                                                             SESSION
FULL ESTIMATED COST
                                                      0.12
                                                              480.55
FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
COPYRIGHT (C) 2008 American Chemical Society (ACS)
Property values tagged with IC are from the ZIC/VINITI data file
provided by InfoChem.
STRUCTURE FILE UPDATES: 11 JUN 2008 HIGHEST RN 1027436-61-8
DICTIONARY FILE UPDATES: 11 JUN 2008 HIGHEST RN 1027436-61-8
```

TSCA INFORMATION NOW CURRENT THROUGH January 9, 2008.

Please note that search-term pricing does apply when

New CAS Information Use Policies, enter HELP USAGETERMS for details.

conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

=> str

:gra c2,dis sia

C~^ C 1 2

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 2

STEREO ATTRIBUTES: NONE :nod 1 g1,2 cb :dis sia

G1 ~ Cb 1 2

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 2

STEREO ATTRIBUTES: NONE :eco 2 e6 c,dis sia

G1 ~ Cb 1 2

NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E6 C AT 2

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 2

STEREO ATTRIBUTES: NONE :gra r66,r665 :dis sia

NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED
ECOUNT IS E6 C AT 2

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

ibon r 5 9 se,r 3 4 m,r 15 19 se,r13 14 n,r 20 24 se,dis sia BOND TYPE/VALUE NOT VALID
The bond type and/or value is not valid. Enter "HELP BOND" for more information.
ibon r 5 9 se,r 3 4 n,r 15 19 se,r13 14 n,r 20 24 se,dis sia NODE AND TYPE NOT VALID
Both a node number and a bond type are not valid. Enter "DIS" to look at the structure. Enter "HELP BOND" for more information.
ibon r 5 9 se,r 3 4 n,r 15 19 se,r 13 14 n,r 20 24 se,dis sia

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED ECOUNT IS E6 C AT 2 GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE

:nod 24 o,dis sia

NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED ECOUNT IS E6 C AT 2

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE :var g1=11/12/22/25 :dis sia

VAR G1=11/12/22/25 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED ECOUNT IS E6 C AT 2

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 25

STEREO ATTRIBUTES: NONE :end L37 STRUCTURE CREATED

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

```
FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
1.2
               TRA L1 1- RN : 16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
            16 SEA L2
L4
            10 L3 AND >=2 46.150.18/RID
L5
             1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
L7
               STR
L8
            29 L7
L9
               SCR 1839
L10
            29 L7 AND L9
L11
               STR L7
L12
       336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
            50 L11 SAM SUB=L12
L14
               STR L11
L15
            32 L14 SAM SUB=L12
               E FURAN/CN
L16
             1 E3
               E THF/CN
1.17
             1 E3
L18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
             2 L22 AND L23 AND L21
L25
            36 L22 AND L23 AND L21 SAM SUB=L18
L26
               STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
            94 L26 AND L22 AND L23 FULL SUB=L18
               SAV TEM G761C1/A L28
L29
               STR L7
L30
             2 L29
L31
               STR L29
L32
             0 L31
L33
               SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L34
             0 L31 AND L33
L35
             33 L31 AND L33 FULL
               SAV TEM G761C1N/A L35
L36
             0 L28, L35 AND L3
    FILE 'STNGUIDE' ENTERED AT 15:14:39 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
L37
               STR
=> 137
SAMPLE SEARCH INITIATED 15:18:40 FILE 'REGISTRY'
```

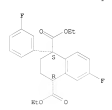
6.2% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

SAMPLE SCREEN SEARCH COMPLETED - 32405 TO ITERATE

```
FULL FILE PROJECTIONS: ONLINE **COMPLETE**
                       BATCH **COMPLETE**
PROJECTED ITERATIONS:
                           637336 TO 658864
PROJECTED ANSWERS:
                            13269 TO 16543
L38
            46 SEA SSS SAM L37
=> 46.150.18/rid and (c5-c6-c6 or c6-c6)/es
      22121069 46.150.18/RID
        155314 C5-C6-C6/ES
       1122454 C6-C6/ES
1.39
       694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
1.2
                TRA I.1 1- RN :
                                    16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
             1 L3 AND OC4-C6-C6/ES
L5
             1 L3 AND C6-C6/ES
L6
L7
               STR
L8
             29 L7
L9
               SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
L12
         336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB=L12
L14
                STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
              1 E3
1.18
         130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
                STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
                STR L20
L22
                SCR 1707 OR 1708
L23
                SCR 1840
L24
             2 L22 AND L23 AND L21
             36 L22 AND L23 AND L21 SAM SUB=L18
L25
L26
                STR L21
1.27
             4 L26 AND L22 AND L23 SAM SUB=L18
             94 L26 AND L22 AND L23 FULL SUB=L18
L28
                SAV TEM G761C1/A L28
```

```
1.29
                STR L7
1.30
              2 L29
L31
                STR L29
              0 L31
                SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L33
L34
              0 L31 AND L33
             33 L31 AND L33 FULL
L35
                SAV TEM G761C1N/A L35
L36
              0 L28, L35 AND L3
     FILE 'STNGUIDE' ENTERED AT 15:14:39 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
1.37
               STR
1,38
             46 L37
L39
        694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES
=> 137 sub=139 sam
SAMPLE SUBSET SEARCH INITIATED 15:19:18 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED -
                                           3321 TO ITERATE
60.2% PROCESSED
                     2000 ITERATIONS
                                                                50 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
PROJECTIONS (WITHIN SPECIFIED SUBSET):
                                                ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET):
                                                    62964 TO 69876
PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET):
                                                        3975 TO
                                                                    5855
L40
             50 SEA SUB=L39 SSS SAM L37
=> d sca
L40 50 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN
    1,4-Naphthalenedicarboxylic acid, 6-fluoro-1-(3-fluorophenyl)-1,2,3,4-
     tetrahydro-, 1,4-diethyl ester, (1R,4S)-rel-
```

MF C22 H22 F2 O4
Relative stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

```
=> d his
     (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)
     FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008
               SET AUTOSEARCH ON
L1
              1 US20050249857/PN
     FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
     FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008
                TRA L1 1- RN :
                                    16 TERMS
    FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008
L3
             16 SEA L2
L4
             10 L3 AND >=2 46.150.18/RID
L5
              1 L3 AND OC4-C6-C6/ES
L6
             1 L3 AND C6-C6/ES
L7
               STR
L8
             29 L7
L9
               SCR 1839
L10
             29 L7 AND L9
               STR L7
         336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB=L12
L14
                STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
              1 E3
L16
               E THF/CN
L17
              1 E3
L18
         130953 L12 AND 16.138.1/RID
L19
             49 L14 SAM SUB=L18
L20
                STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
                STR L20
L22
                SCR 1707 OR 1708
L23
               SCR 1840
L24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
             94 L26 AND L22 AND L23 FULL SUB=L18
                SAV TEM G761C1/A L28
L29
                STR L7
L30
             2 L29
L31
                STR L29
L32
             0 L31
L33
                SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L34
             0 L31 AND L33
L35
             33 L31 AND L33 FULL
                SAV TEM G761C1N/A L35
1.36
             0 L28, L35 AND L3
```

FILE 'STNGUIDE' ENTERED AT 15:14:39 ON 12 JUN 2008

FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008

1.37 STR

L38 46 L37

L39 694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES L40 50 L37 SAM SUB=L39

=> str 137 :dis

VAR G1=11/12/22/25 :att o 23 vn d,dis sia

1 2

0 26 19 23

VAR G1=11/12/22/25 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED ECOUNT IS E6 C AT 2

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 26

STEREO ATTRIBUTES: NONE :end L41 STRUCTURE CREATED

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 TRA L1 1- RN : 16 TERMS

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 L3 16 SEA L2

L4 10 L3 AND >=2 46.150.18/RID 1.5

L6 L7

L8

L9

L10

L11

1 L3 AND OC4-C6-C6/ES

1 L3 AND C6-C6/ES

STR

SCR 1839

29 L7 AND L9

STR L7

29 L7

```
336455 >=2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L12
L13
            50 L11 SAM SUB=L12
L14
               STR L11
L15
            32 L14 SAM SUB=L12
               E FURAN/CN
L16
             1 E3
               E THF/CN
1.17
             1 E3
L18
        130953 L12 AND 16.138.1/RID
L19
            49 L14 SAM SUB=L18
L20
               STR L14
    FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
               STR L20
L22
               SCR 1707 OR 1708
L23
               SCR 1840
L24
             2 L22 AND L23 AND L21
L25
            36 L22 AND L23 AND L21 SAM SUB=L18
L26
               STR L21
L27
             4 L26 AND L22 AND L23 SAM SUB=L18
L28
            94 L26 AND L22 AND L23 FULL SUB=L18
               SAV TEM G761C1/A L28
L29
               STR L7
L30
             2 L29
L31
               STR L29
L32
             0 L31
L33
               SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L34
             0 L31 AND L33
L35
             33 L31 AND L33 FULL
               SAV TEM G761C1N/A L35
L36
             0 L28, L35 AND L3
    FILE 'STNGUIDE' ENTERED AT 15:14:39 ON 12 JUN 2008
    FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
1.37
               STR
L38
            46 L37
L39
        694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES
L40
             50 L37 SAM SUB=L39
L41
               STR L37
=> 141 sub=139 sam
SAMPLE SUBSET SEARCH INITIATED 15:20:22 FILE 'REGISTRY'
SAMPLE SUBSET SCREEN SEARCH COMPLETED - 3321 TO ITERATE
60.2% PROCESSED
                   2000 ITERATIONS
                                                             50 ANSWERS
INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)
SEARCH TIME: 00.00.01
PROJECTIONS (WITHIN SPECIFIED SUBSET): ONLINE **COMPLETE**
PROJECTED ITERATIONS (WITHIN SPECIFIED SUBSET): 62964 TO 69876
```

PROJECTED ANSWERS (WITHIN SPECIFIED SUBSET):

3975 TO 5855

L42 50 SEA SUB=L39 SSS SAM L41

=> d sca

L42 50 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN 1-Naphthalenol, 4-(2,4-dimethylphenyl)-3-ethenyl-1,2,3,4-tetrahydro-6-methoxy-2-methyl-, (1R,2R,3R,4S)-rel-

F C22 H26 O2

Relative stereochemistry.

PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):.

L42 50 ANSWERS REGISTRY COPYRIGHT 2008 ACS on STN

IN Glycine, N-[(4-methylphenyl)sulfonyl]-N-[(1R,2R)-1,2,3,4-tetrahydro-7-1]

methoxy-1-phenyl-2-naphthalenyl]-

MF C26 H27 N O5 S

Absolute stereochemistry.

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):0

=> 141 sub=139 full

FULL SUBSET SEARCH INITIATED 15:20:48 FILE 'REGISTRY'
FULL SUBSET SCREEN SEARCH COMPLETED - 67200 TO ITERATE

100.0% PROCESSED 67200 ITERATIONS

7953 ANSWERS

SEARCH TIME: 00.00.01

L43 7953 SEA SUB=L39 SSS FUL L41

=> sav tem g761c1n2/a 143

=> log h

 COST ÎN U.S. DOLLARS
 SINCE FILE
 TOTAL

 ENTRY
 SESSION

 FULL ESTIMATED COST
 197.49
 678.04

SESSION WILL BE HELD FOR 120 MINUTES

STN INTERNATIONAL SESSION SUSPENDED AT 15:21:07 ON 12 JUN 2008

Connecting via Winsock to STN

Welcome to STN International! Enter x:X

LOGINID:SSPTASYG1600

PASSWORD:

* * * * * * RECONNECTED TO STN INTERNATIONAL * * * * * * SESSION RESUMED IN FILE 'REGISTRY' AT 15:23:35 ON 12 JUN 2008

FILE 'REGISTRY' ENTERED AT 15:23:35 ON 12 JUN 2008 COPYRIGHT (C) 2008 American Chemical Society (ACS)

COST IN U.S. DOLLARS SINCE FILE ENTRY

FULL ESTIMATED COST

ENTRY SESSION 197.49 678.04

TOTAL

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON

L1 1 US20050249857/PN

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008

FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008

L3 16 SEA L2

L4 10 L3 AND >=2 46.150.18/RID

L5 1 L3 AND OC4-C6-C6/ES

L6 1 L3 AND C6-C6/ES

```
STR
1.8
             29 L7
L9
                SCR 1839
L10
             29 L7 AND L9
L11
               STR L7
L12
        336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES
L13
             50 L11 SAM SUB=L12
L14
                STR L11
L15
             32 L14 SAM SUB=L12
               E FURAN/CN
L16
              1 E3
               E THF/CN
              1 E3
L18
         130953 L12 AND 16.138.1/RID
L19
             49 L14 SAM SUB=L18
L20
                STR L14
     FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008
L21
                STR L20
L22
                SCR 1707 OR 1708
                SCR 1840
1.24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
L27
              4 L26 AND L22 AND L23 SAM SUB=L18
L28
             94 L26 AND L22 AND L23 FULL SUB=L18
                SAV TEM G761C1/A L28
L29
                STR L7
L30
             2 L29
L31
                STR L29
L32
              0 L31
L33
                SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L34
              0 L31 AND L33
L35
             33 L31 AND L33 FULL
                SAV TEM G761C1N/A L35
L36
              0 L28, L35 AND L3
     FILE 'STNGUIDE' ENTERED AT 15:14:39 ON 12 JUN 2008
     FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
L37
                STR
1.38
             46 L37
1.39
         694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES
L40
             50 L37 SAM SUB=L39
L41
                STR L37
L42
             50 L41 SAM SUB=L39
L43
           7953 L41 FULL SUB=L39
                SAV TEM G761C1N2/A L43
=> fil caplus
COST IN U.S. DOLLARS
                                                  SINCE FILE
                                                       ENTRY
                                                               SESSION
FULL ESTIMATED COST
                                                      197.95
                                                                678.50
FILE 'CAPLUS' ENTERED AT 15:24:14 ON 12 JUN 2008
```

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=> s 128

L44 86 L28

=> d hitstr 1-86

L44 ANSWER 1 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 957467-24-2P

RL: ANT (Analyte); PRP (Properties); PUR (Purification or recovery); ANST (Analytical study); PREP (Preparation)

(new lignans from the heartwood of Cunninghamia lanceolata)

RN 957467-24-2 CAPLUS

CN 2(3H)-Furanone, 3-(acetyloxy)-3-[[4-(acetyloxy)-3,5-

dimethoxyphenyl]methyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, (3S,4S)- (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L44 ANSWER 2 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

T 947685-66-7

RL: BSU (Biological study, unclassified); BIOL (Biological study) (Content of lignin; structure of lignins in developing xylem of Norway spruce (Picea abies))

RN 947685-66-7 CAPLUS

CN Phenol, 4,4'-[(tetrahydro-3,4-furandiyl)bis(methylene)]bis[2-methoxy-,
1,1'-diacetate (CA INDEX NAME)

L44 ANSWER 3 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 54797-70-5P, Matairesinol diacetate 578013-40-8P, Matairesinol dibutyrate 578014-11-6P, Matairesinol distearate 578014-47-8P, Matairesinol disuccinate 578014-71-8P, Matairesinol bis(methyl succinate) 578017-11-5P

578017-13-7P

RL: FFD (Food or feed use); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of lignan ester derivs. for use in pharmaceutical compns. and as dietary supplements)

RN 54797-70-5 CAPLUS

2(3H)-Furanone, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, (3R,4R)- (CA INDEX NAME)

Absolute stereochemistry.

RN 578013-40-8 CAPLUS

CN Butanoic acid, [(3R,4R)-tetrahydro-2-oxo-3,4-furandiyl]bis[methylene(2-methoxy-4,1-phenylene)] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 578014-11-6 CAPLUS

CN Octadecanoic acid, [(3R,4R)-tetrahydro-2-oxo-3,4-furandiy1]bis[methylene(2-

methoxy-4,1-phenylene)] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 578014-47-8 CAPLUS

CN Butanedioic acid, [(3R,4R)-tetrahydro-2-oxo-3,4-furandiy1]bis[methylene(2-methoxy-4,1-phenylene)] ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

- CO2H

RN 578014-71-8 CAPLUS

CN Butanedioic acid, [(3R,4R)-tetrahydro-2-oxo-3,4-furandiyl]bis[methylene(2-methoxy-4,1-phenylene)] dimethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-B

- RN 578017-11-5 CAPLUS
- CN Butanoic acid, [(3R,4R)-tetrahydro-2-oxo-3,4-furandiyl]bis(methylene-3,1-phenylene) ester, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

- RN 578017-13-7 CAPLUS
- CN Butanoic acid, 4-[(S)-hydroxy[(3R,4R)-tetrahydro-4-[[3-methoxy-4-(1-oxobutoxy)phenyl]methyl]-5-oxo-3-furanyl]methyl]-2-methoxyphenyl ester (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 4 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 406955-40-6P, Renealtin A acetate
 - RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (renealtins A and B, new diarylheptanoids with THF ring from seeds of Renealmia exaltata)
- RN 406955-40-6 CAPLUS
- CN Ethanone, 2-[(2R,4R,5R)-4-(acetyloxy)-5-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-2-furanyl]-1-[4-(acetyloxy)-3-methoxyphenyl]-, rel-(9CI) (CA INDEX NAME)

Relative stereochemistry.

L44 ANSWER 5 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

56440-75-6

RL: PRP (Properties) (properties of)

56440-75-6 CAPLUS RN

3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3methoxyphenyl]methyl]tetrahydro-4-hydroxy-, α-acetate, (2S,3R,4S)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L44 ANSWER 6 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

211371-46-9, Taxiresinol tetraacetate

RL: BAC (Biological activity or effector, except adverse); BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence)

(bioactive lignans and taxoids from Taxus mairei riits)

211371-46-9 CAPLUS RN

1,2-Benzenediol, 4-[(2S,3R,4R)-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-CN [(acetyloxy)methyl]tetrahydro-2-furanyl]-, diacetate (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

- L44 ANSWER 7 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- 72092-51-4

RL: BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); BIOL (Biological study); PROC (Process)

(preparation of lignans and interaction with sex hormone-binding globulin) RN

72092-51-4 CAPLUS

CN Phenol, 4,4'-[(tetrahydro-3,4-furandiyl)bis(methylene)]bis[2-methoxy-, diacetate, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

L44 ANSWER 8 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 198827-24-6P, 7(R)-Methoxy-8-epi-matairesinol diacetate

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and properties of; bioactive iridoids and a new lignan from Allamanda cathartica and Himatanthus fallax from the Suriname rainforest)

RN 198827-24-6 CAPLUS

CN 2(3H)-Furanone, 3-[[4-(acetyloxy)-3-methoxyphenyl]methoxymethyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, [3R-[3\(\alpha(R^*)\), 4\(\alpha)]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 9 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 191481-13-7P, 5,5'-Dimethoxy-9-0-(β-D-

glucopyranosyl)lariciresinol hexaacetate

RL: BOC (Biological occurrence); BSU (Biological study, unclassified); PRP (Properties); PUR (Purification or recovery); BIOL (Biological study);

OCCU (Occurrence); PREP (Preparation)
(alkaloids and other compds. from seeds of Tabernaemontana cymosa)

RN 191481-13-7 CAPLUS

CN β-D-Glucopyranoside, [(2S,3R,4R)-2-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[(4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]tetrahydro-3-furanyl]methyl tetraacetate (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L44 ANSWER 10 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

T 189204-64-6P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and properties of)

RN 189204-64-6 CAPLUS

CN 2(3H)-Furanone, 4-[(acetyloxy)[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, [3R-[3α, 4β(R*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

- L44 ANSWER 11 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 178178-13-7
 - RL: BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence)

(sesquiterpene lactones and other constituents of Stevia maimarensis and Synedrellopsis grisebachii)

RN 178178-13-7 CAPLUS

CN 3-Furanmethanol, α-[3-(acetyloxy)-4-methoxyphenyl]-4-[[3-(acetyloxy)-4-methoxyphenyl]methyl]tetrahydro-, acetate (9CI) (CA INDEX NAME)

L44 ANSWER 12 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 119740-40-8 158042-34-3 158111-16-1 158189-07-2

RL: BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence) (from Abies marocana wood)

RN 119740-40-8 CAPLUS

CN 2-Propenoic acid, 3-[4-(acetyloxy)phenyl]-, [2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3-furanyl]methyl ester, [2S- $(2\alpha, 3\beta, 4\beta)$]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

RN 158042-34-3 CAPLUS

CN 3-Furanmethanol, 5-(acetyloxy)-2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2S-(2 α , 3 β , 4 β , 5 α)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RN 158111-16-1 CAPLUS

CN 3,4-Furandimethanol, α 4,2-bis[4-(acetyloxy)-3-methoxyphenyl]tetrahydro-, diacetate, [2S-[2 α ,3 β ,4 β (R*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 158189-07-2 CAPLUS

CN 3,4-Furandimethanol, α 4,2-bis[4-(acetyloxy)-3-methoxyphenyl]tetrahydro-, diacetate, [2S-[2 α ,3 β ,4 β (S*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and properties of)

RN 67560-67-2 CAPLUS

CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [28-(2α, 3β, 4β)]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 13 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 166983-32-0P, Busaliol triacetate 166983-33-1P
 RI: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)
 (preparation and NMR data of)

RN 166983-32-0 CAPLUS

CN 3-Furanmethanol, α-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate (9CI) (CA INDEX NAME)

RN 166983-33-1 CAPLUS

CN 3-Furanmethanol, α-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]ethoxymethyl]tetrahydro-, acetate (9CI) (CA INDEX NAME)

L44 ANSWER 14 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 154634-44-3

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(biol. activity of secondary metabolites from Bupleurum salicifolium (Umbelliferae))

RN 154634-44-3 CAPLUS

CN 2(3H)-Furanone, 3-(acetyloxy)-3,4-bis[[4-(acetyloxy)-3-

methoxyphenyl]methyl]dihydro-, (3S-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 15 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- TT 56440 75
 - RL: BOC (Biological occurrence); BSU (Biological study, unclassified); BIOL (Biological study); OCCU (Occurrence)
 - (chemical study on the genus Prunus (Rosaceae): comparative studies on the chemical constituents of the barks of the subgenera Cerasus, Padus and
- Laurocerasus plants) RN 56440-75-6 CAPLUS
- CN 3-Furanmethanol, 2-(4-(acetyloxy)-3-methoxyphenyl)-4-[(4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-4-hydroxy-, α-acetate, (2S,3R,4S)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

L44 ANSWER 16 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 159821-58-6

RL: ANT (Analyte); ANST (Analytical study) (determination of phenolic extractives from wood of Salix sachalinensis Fr. Schm.)

RN 159821-58-6 CAPLUS

CN Benzoic acid, 4-(acetyloxy)-, [4-(acetyloxy)[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]-2-[4-(acetyloxy)-3,5-dimethoxyphenyl]tetrahydro-3-furanyl]methyl ester (CA INDEX NAME)

- L44 ANSWER 17 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 158372-31-7

RL: ANT (Analyte); BOC (Biological occurrence); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); OCCU (Occurrence)

(lignans of Abies koreana)

RN 158372-31-7 CAPLUS CN 3-Furanmethanol α-

3-Furanmethanol, α -[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-5-methoxy-, acetate (9CI) (CA INDEX NAME)

L44 ANSWER 18 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
11 124265-87-8 18111-17-2
 RL: BIOL (Biological study)
 (from acetylated Abies pinsapo wood)
RN 124265-87-8 CAPLUS
CN 3-Furanmethanol, 5-(acetyloxy)-α-[4-(acetyloxy)-3-methoxyphenyl]-4[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate,

 $[3R-[3\alpha(S^*),4\beta,5\alpha]]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

RN 158111-17-2 CAPLUS
CN 2-Propenoic acid, 3-[4-(acetyloxy)phenyl]-, [2-[4-(acetyloxy)-3methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3furanyl]methyl ester, [2S-[2a,3β(E),4β]]- (9CI) (CA INDEX
NAME)

Absolute stereochemistry.
Double bond geometry as shown.

IT 158042-34-3P 158042-36-5P 158111-16-1P 158189-07-2P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and spectral properties of)

- RN 158042-34-3 CAPLUS
- CN 3-Furanmethanol, 5-(acetyloxy)-2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [25-(2a,3b,4b,5a)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

- RN 158042-36-5 CAPLUS
- CN 2-Furanol, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, acetate, [2S-(2\alpha,3\beta,4\alpha)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 158111-16-1 CAPLUS

CN 3,4-Furandimethanol, $\alpha 4$,2-bis[4-(acetyloxy)-3-methoxyphenyl]betrahydro-, diacetate, [2S-[2 α ,3 β ,4 β (R*)]]-(9C1) (CA INDEX NAME)

Absolute stereochemistry.

RN 158189-07-2 CAPLUS

CN 3,4-Furandimethanol, α 4,2-bis[4-(acetyloxy)-3-methoxypheny]|tetrahydro-, diacetate, [2S-[2 α ,3 β ,4 β (S*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 19 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 156616-60-3P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation)

(preparation and NMR of)

RN 156616-60-3 CAPLUS

CN 2(3H)-Furanone, 3-(acetyloxy)-3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, (3R-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 20 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 154461-66-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 154461-66-2 CAPLUS

CN 3-Furanmethanol, 2-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]tetrahydro-, acetate, (2\alpha,3\beta,4\beta)- (9CI) (CA INDEX NAME)

Relative stereochemistry.

- L44 ANSWER 21 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 154634-44-3
 RL: BIOL (Biological study)

(potato cyst nematode hatch inhibition by, from Bupleurum salicifolium)

RN 154634-44-3 CAPLUS

CN 2(3H)-Furanone, 3-(acetyloxy)-3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, (3S-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- RN 67560-67-2 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2S-(2α, 3β, 4β)]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 23 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 136051-42-8P
 - RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)
- RN 136051-42-8 CAPLUS
- CN 3-Furanmethanol, 4-[[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]-2-[4-(acetyloxy)-3-methoxyphenyl]tetrahydro-, acetate, [2S-(2\alpha, 3\beta, 4\beta)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 24 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 136051-42-8P
 - RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)
- RN 136051-42-8 CAPLUS
- CN 3-Furanmethanol, 4-[[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]-2-[4-(acetyloxy)-3-methoxyphenyl]tetrahydro-, acetate, [2S-(2a,3B,4B)]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 25 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 96087-12-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and oxidation of)

RN 96087-12-6 CAPLUS

CN 3,4-Furandiol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-[(acetyloxy)methyl]tetrahydro-, [2R-(2a, 3a, 4a)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- IT 56440-75-6P 133137-65-2P
 RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)
- RN 56440-75-6 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-4-hydroxy-, α-acetate, (2S, 3R, 4S)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

RN 133137-65-2 CAPLUS

CN Methanone, $[4-(acetyloxy)-3-methoxyphenyl][5-[4-(acetyloxy)-3-methoxyphenyl]-4-[(acetyloxy)methyl]tetrahydro-3-furanyl]-, [3R-<math>(3\alpha, 4\alpha, 5\beta)$]- (9C1) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 26 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 131653-23-1P 131723-85-8P
 RL: SPN (Synthetic preparation); PREP (Preparation)
- (preparation of)
- RN 131653-23-1 CAPLUS CN β-D-Glucopyranoside,
 - ZN B-D-Glucopyranoside, [4-(acetyloxy)-3-methoxyphenyl] [5-[4-(acetyloxy)-3-methoxyphenyl]-4-[(acetyloxy)methyl)tetrahydro-3-furanyl)methyl, tetraacetate, [38-[3α(R*), 4β, 5α]]- (9CI) (CA INDEX NAME)

- RN 131723-85-8 CAPLUS
- CN β -D-Allopyranoside, [4-(acetyloxy)-3-methoxyphenyl][5-[4-(acetyloxy)-3-methoxyphenyl]-4-[(acetyloxy)methyl]tetrahydro-3-furanyl]methyl, tetraacetate, [3S- $3\alpha(R^4)$, 4β , 5α]- [9C1) (CA INDEX NAME)

L44 ANSWER 27 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 126906-02-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of (Erratum))

RN 126906-02-3 CAPLUS

CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3-hydroxy-, α-acetate, [2R-(2α, 3α, 4β)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

- L44 ANSWER 28 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 126906-02-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

- RN 126906-02-3 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3-hydroxy-, α-acetate, [2R-(2α, 3α, 4β)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

L44 ANSWER 29 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 126882-64-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 126882-64-2 CAPLUS

CN β-D-Xylopyranoside, [2-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]tetrahydro-3-furanyl]methyl, triacetate, [25-(2α,3β,4β)]- (9C1) (CA INDEX NAME)

L44 ANSWER 30 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 126026-27-5P RL: SPN (Synthetic

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 126026-27-5 CAPLUS

CN Phenol, 4-[4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3-furanyl]-2-methoxy-, acetate, (3S-trans)- (9CI) (CA INDEX NAME)

L44 ANSWER 31 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN ΙT 124265-87-8P 124265-88-9P

RL: PREP (Preparation)

(from firwood, structure of) 124265-87-8 CAPLUS

RN

3-Furanmethanol, 5-(acetyloxy)- α -[4-(acetyloxy)-3-methoxyphenyl]-4-[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, CN $[3R-[3\alpha(S^*), 4\beta, 5\alpha]]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

RN 124265-88-9 CAPLUS

3-Furanmethano1, 5-(acetyloxy)- α -[4-(acetyloxy)-3-methoxyphenyl]-4-[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, CN $[3R-[3\alpha(R^*), 4\beta, 5\alpha]]-(9CI)$ (CA INDEX NAME)

L44 ANSWER 32 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 122585-41-5P, Lophirochalcone undecaacetate 122621-93-6P , Isombamichalcone hexaacetate

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 122585-41-5 CAPLUS

CN 2-Propen-1-one, 3-[3-[[2,4-bis(acetyloxy)-5-[5-[4-(acetyloxy)phenyl]-3-[[4-(acetyloxy)phenyl]methyl]-4-[2,4-bis(acetyloxy)benzoyl]tetrahydro-2-furanyl]phenyl][2,4-bis(acetyloxy)phenyl]methyl]-2-[4-(acetyloxy)phenyl]-2,3-dihydro-5-benzofuranyl]-1-[2,4-bis(acetyloxy)phenyl]-, [2 α [R*[2R*,3R*,5(E)]],3 α ,4 β ,5 α]- (9CI) (CA INDEX NAME)

Relative stereochemistry.
Double bond geometry as shown.
Currently available stereo shown.

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RN 122621-93-6 CAPLUS

CN Methanone, [2-[4-(acetyloxy)phenyl]-4-[[4-(acetyloxy)phenyl]methyl]-5-[2,4-bis(acetyloxy)phenyl]tetrahydro-3-furanyl][2,4-bis(acetyloxy)phenyl]-, (2a,3B,4a,5a) - (9CI) (CA INDEX NAME)

Relative stereochemistry. Currently available stereo shown.

- L44 ANSWER 33 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
 - T 67560-67-2P 77255-60-8P 119740-40-8P RL: SPN (Synthetic preparation); PREP (Preparation)
- (preparation of) RN 67560-67-2 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[(4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2S-(2 α , 3 β , 4 β)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 77255-60-8 CAPLUS

CN 3,4-Furandimethanol, $\alpha 4$,2-bis[4-(acetyloxy)-3methoxyphenyl]tetrahydro-, diacetate (9CI) (CA INDEX NAME)

RN 119740-40-8 CAPLUS

CN 2-Propenoic acid, 3-[4-(acetyloxy)phenyl]-, [2-[4-(acetyloxy)-3methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3furanyl]methyl ester, $[2S-(2\alpha, 3\beta, 4\beta)]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

L44 ANSWER 34 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

119264-66-3P, Mbamichalcone hexaacetate

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 119264-66-3 CAPLUS

CN Methanone, [2-[4-(acetyloxy)phenyl]-4-[[4-(acetyloxy)phenyl]methyl]-5-[2,4-bis(acetyloxy)phenyl]tetrahydro-3-furanyl][2,4-bis(acetyloxy)phenyl]-, (2a, 3B, 4a, 5B)- (9CI) (CA INDEX NAME)

Relative stereochemistry.

Currently available stereo shown.

L44 ANSWER 35 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 117458-40-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 117458-40-9 CAPLUS

CN Methanone, [2,5-bis[4-(acetyloxy)phenyl]tetrahydro-3,4-furandiyl]bis[[2,4-bis(acetyloxy)phenyl]- (9CI) (CA INDEX NAME)

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L44 ANSWER 36 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 116384-20-4P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 116384-20-4 CAPLUS

CN 3-Furanmethanol, 2-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]tetrahydro-, acetate, [2S-(2\alpha,3\beta,4\beta)] (OCI) (CA INDEX NAME)

L44 ANSWER 37 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 107783-49-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and deacetylation of)

RN 107783-49-3 CAPLUS

CN 3-Furanmethanol, 4-(acetyloxy)-2-[4-(acetyloxy)-3-methoxyphenyl]-4-[{4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2S-(2a, 3B, 4a)]- (9CI) (CA INDEX NAME)

L44 ANSWER 38 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 107783-49-3P, Olivil tetraacetate

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 107783-49-3 CAPLUS

CN 3-Furanmethanol, 4-(acetyloxy)-2-[4-(acetyloxy)-3-methoxyphenyl]-4-[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2S-(2a, 3B, 4a)]- (9CI) (CA INDEX NAME)

- L44 ANSWER 39 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 106647-16-9P RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 106647-16-9 CAPLUS

Note: 10008;1-9 CARLDON
2(3H)-Furanone, 4-[(4-(acetyloxy)-3-methoxyphenyl]methyl]-3-[(4-(acetyloxy)-3-methoxy-5-(2,3,4,6-tetra-0-acetyl-B-D-glucopyranosyl)phenyl]methyl]dihydron, (3R-trans)- (9CI) (CA INDEX NAME)

L44 ANSWER 40 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 105233-16-7P 105308-08-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

N 105233-16-7 CAPLUS

CN 3-Furanmethanol, 2-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, $(2\alpha,3\beta,4\beta)$ - (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 105308-08-5 CAPLUS CN 3-Furanmethanol, 2-

3-Furanmethanol, $2-[4-(acetyloxy)-3-methoxypheny1]-4-[[4-(acetyloxy)-3-methoxypheny1]methy1]tetrahydro-, acetate, <math>(2\alpha, 3\beta, 4\beta)-(9C1)$ (CA INDEX NAME)

L44 ANSWER 41 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 104086-79-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 104086-79-5 CAPLUS

CN 3-Furanmethanol, α-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate (9CI) (CA INDEX NAME)

L44 ANSWER 42 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 101218-38-6P 101218-39-7P 101247-19-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 101218-38-6 CAPLUS

CN Phenol, $4-[[4-(acetyloxy)-3-methoxyphenyl]methoxymethyl]tetrahydro-2-methoxy-furanyl]methyl]-2-methoxy-, acetate, [2R-[2<math>\alpha$, 3 β , 4α (8*)]]- (9C1) (CA INDEX NAME)

RN 101218-39-7 CAPLUS

CN Phenol, $4-[[4-(acetyloxy)-3-methoxyphenyl]methoxymethyl]tetrahydro-2-methoxy-3-furanyl]methyl]-2-methoxy-, acetate, [2S-[2<math>\alpha$, 3 α , 4 β (R*)]]- [9CI) (CA INDEX NAME)

RN 101247-19-2 CAPLUS

CN Phenol, 4-[[4-[[4-(acetyloxy)-3-methoxyphenyl]methoxymethyl]tetrahydro-2-methoxy-3-furanyl]methyl]-2-methoxy-, acetate, [2R-[2a, 3B, 4a(R*)]]- [9CI) (CA INDEX NAME)

L44 ANSWER 43 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 98770-68-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 98770-68-4 CAPLUS

CN 2(3H)-Furanone, 4-[4-(acetyloxy)-3-methoxybenzoy1]-3-[[4-(acetyloxy)-3methoxyphenyl]methyl]dihydro-, (3R-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 44 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

ΙT 57934-45-9

RL: PRP (Properties) (proton NMR of)

RN 57934-45-9 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L44 ANSWER 45 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 96917-10-1P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN

96917-10-1 CAPLUS

2-Propenoic acid, 3-[4-(acetyloxy)-3-methoxyphenyl]-, 4-[[4-[[4-CN (acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3-furanyl]methyl]-2methoxyphenyl ester, trans-(-)- (9CI) (CA INDEX NAME)

Rotation (-). Absolute stereochemistry unknown. Double bond geometry unknown.

- L44 ANSWER 46 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 96087-12-6P

RL: SPN (Synthetic preparation); PREP (Preparation)

- (preparation of) RN 96087-12-6 CAPLUS
- CN 3,4-Furandiol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-[(acetyloxy)methyl]tetrahydro-, [2R-(2α, 3α, 4α)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 47 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 21497-66-5 57934-45-9 RL: PRP (Properties)
 - (electronic spectrum of)
- RN 21497-66-5 CAPLUS
- CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 57934-45-9 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

- L44 ANSWER 48 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- II 83327-17-7P
 RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)
- RN 83327-17-7 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2R-(2α , 3 β , 4 β)]-(9CI) (CA INDEX NAME)

L44 ANSWER 49 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 81262-98-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 81262-98-8 CAPLUS

CN 3-Furanmethanol, 5-(acetyloxy)-α-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate (9CI) (CA INDEX NAME)

L44 ANSWER 50 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 81613-44-7P

CN

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 81613-44-7 CAPLUS

 β -D-Glucopyranoside, [2,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3-furanyl]methyl, tetraacetate (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 51 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

ΙT 78032-16-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 78032-16-3 CAPLUS

CN

2(3H)-Furanone, 3,4-bis[[3-(acetyloxy)phenyl]methyl]dihydro-, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

- L44 ANSWER 52 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- ΙT 79863-74-4 79863-75-5 RL: BIOL (Biological study)

(in compression wood of larch)

RN 79863-74-4 CAPLUS CN

2(3H)-Furanone, 3-[[4-(acetyloxy)-3-[2-(acetyloxy)-1-[[4-(acetyloxy)-3methoxyphenyl]methyl]ethyl]-5-methoxyphenyl]methyl]-4-[[4-(acetyloxy)-3methoxyphenyl]methyl]dihydro- (CA INDEX NAME)

RN 79863-75-5 CAPLUS

CN 2(3H)-Furanone, 4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-[[2',6-bis(acetyloxy)-5'-[3-(acetyloxy)propyl]-3',5-dimethoxy[1,1'-biphenyl]-3-yl]methyl]dihydro- (9CI) (CA INDEX NAME)

- L44 ANSWER 53 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 67560-67-2P 77225-34-4P 77255-60-8P
 - RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)
- RN 67560-67-2 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2S- $(2\alpha, 3\beta, 4\beta)$]-(9C1) (CA INDEX NAME)

Absolute stereochemistry.

- RN 77225-34-4 CAPLUS
- CN Methanone, [4-(acetyloxy)-3-methoxyphenyl][5-[4-(acetyloxy)-3-methoxyphenyl]-4-[(acetyloxy)methyl]tetrahydro-3-furanyl]- (CA INDEX NAME)

RN 77255-60-8 CAPLUS

OMe

CN 3,4-Furandimethanol, α4,2-bis[4-(acetyloxy)-3-methoxyphenyl]tetrahydro-, diacetate (9CI) (CA INDEX NAME)

- 78032-16-3P RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
 - (Reactant or reagent) (preparation and hydrolysis of)
 - 78032-16-3 CAPLUS RN
 - 2(3H)-Furanone, 3,4-bis[[3-(acetyloxy)phenyl]methyl]dihydro-, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

- ΙT 78032-15-2P
 - RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
- (preparation and reduction of) RN 78032-15-2 CAPLUS
- CN
- 2,5-Furandione, 3,4-bis[[3-(acetyloxy)phenyl]methyl]dihydro-, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

- L44 ANSWER 55 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- ΙT 75679-27-5
 - RL: PRP (Properties) (NMR spectrum of)
- RN 75679-27-5 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]tetrahydro-, acetate (9CI) (CA INDEX NAME)

IT 67560-67-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 67560-67-2 CAPLUS

OAc OMe

CN 3-Furanmethanol, $2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, <math>[2S-(2\alpha,3\beta,4\beta)]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 57 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 67560-67-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

- RN 67560-67-2 CAPLUS
- CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, $[2S-(2\alpha, 3\beta, 4\beta)]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 58 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

ΙT 55665-05-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN

55665-05-9 CAPLUS
Methanone, [4-(acetyloxy)-3,5-dimethoxyphenyl][5-[4-(acetyloxy)-3,5dimethoxyphenyl]-4-[(acetyloxy)methyl]tetrahydro-3-furanyl]- (CA INDEX NAME)

- L44 ANSWER 59 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- ΙT 72448-84-1P 72448-85-2P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

72448-84-1 CAPLUS RN

CN 2-Propenoic acid, 3-[4-(acetyloxy)phenyl]-, [2-[4-(acetyloxy)-3methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3furanyl]methyl ester (CA INDEX NAME)

72448-85-2 CAPLUS 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)phenyl]methyl]tetrahydro-, acetate (9CI) (CA INDEX NAME)

L44 ANSWER 60 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN ΙT 72092-51-4

RL: PRP (Properties) (NMR spectrum of, as model for lignin)

RN 72092-51-4 CAPLUS

CN Phenol, 4,4'-[(tetrahydro-3,4-furandiyl)bis(methylene)]bis[2-methoxy-, diacetate, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

L44 ANSWER 61 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 69394-07-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 69394-07-6 CAPLUS

CN 2(3H)-Furanone, 3,4-bis[[4-(acetyloxy)-3-[2-(acetyloxy)-2-[4-(acetyloxy)-3methoxyphenyl]-1-[(acetyloxy)methyl]ethyl]-5-methoxyphenyl]methyl]dihydro-(9CI) (CA INDEX NAME)

PAGE 2-A

L44 ANSWER 62 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 67560-67-2P

RL: PRP (Properties); SPN (Synthetic preparation); PREP (Preparation) (preparation and carbon-13 NMR of)

RN 67560-67-2 CAPLUS

CN 3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-, acetate, [2S-(2α , 3β , 4β)]-(9C1) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 63 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

67308-37-6P ΙT

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 67308-37-6 CAPLUS

β-D-Glucopyranoside, [2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]tetrahydro-3-furanyl]methyl, tetraacetate, (2α,3β,4β)-(-)- (9CI) (CA INDEX NAME)

- L44 ANSWER 64 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- ΙT 55665-05-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN

55665-05-9 CAPLUS
Methanone, [4-(acetyloxy)-3,5-dimethoxyphenyl][5-[4-(acetyloxy)-3,5-CN dimethoxyphenyl]-4-[(acetyloxy)methyl]tetrahydro-3-furanyl]- (CA INDEX NAME)

L44 ANSWER 65 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 65560-00-1P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 65560-00-1 CAPLUS

CN 1,2-Benzenediol, 4-[(acetyloxy)[4-(acetyloxy)-3-[4-(acetyloxy)phenyl]tetrahydro-2-furanyl]methyl]-, diacetate,

 $[2S-[2\alpha(R^*),3\beta,4\alpha]]-(9CI)$ (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 66 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN IT 21497-65-4 21497-66-5
- IT 21497-65-4 21497-66-RL: PRP (Properties)

(UV spectrum of) RN 21497-65-4 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (Z,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 21497-66-5 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 57934-45-9

RL: RCT (Reactant); RACT (Reactant or reagent) (photochem. rearrangement of)

RN 57934-45-9 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

- L44 ANSWER 67 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 64855-03-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 64855-03-4 CAPLUS

CN 2(3H)-Furanone, 4-[[4-(acetyloxy)-3-[2-(acetyloxy)-1-[(acetyloxy) [4-(acetyloxy)-3-methoxyphenyl]methyl]ethyl]-5-methoxyphenyl]methyl]inethyl

L44 ANSWER 68 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 63339-52-6

RL: PROC (Process)

(photochromism of, reaction mechanism of)

RN 63339-52-6 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-(CA INDEX NAME)

L44 ANSWER 69 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 61504-11-8P 61504-12-9P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 61504-11-8 CAPLUS

CN 2(3H)-Furanone, 3-(acetyloxy)-3,4-bis[[4-(acetyloxy)-3-

methoxyphenyl]methyl]dihydro-, (3R-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 61504-12-9 CAPLUS

CN 2(3H)-Furanone, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-3hydroxy-, (3R-cis)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 70 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 77255-60-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 77255-60-8 CAPLUS

CN 3,4-Furandimethanol, α4,2-bis[4-(acetyloxy)-3methoxyphenyl]tetrahydro-, diacetate (9CI) (CA INDEX NAME)

L44 ANSWER 71 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 57934-45-9

RL: RCT (Reactant); RACT (Reactant or reagent)

(photolysis of, mechanism of)

RN 57934-45-9 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L44 ANSWER 72 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

56440-75-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

56440-75-6 CAPLUS

3-Furanmethanol, 2-[4-(acetyloxy)-3-methoxyphenyl]-4-[[4-(acetyloxy)-3methoxyphenyl]methyl]tetrahydro-4-hydroxy-, α-acetate, (2S, 3R, 4S)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (-).

- L44 ANSWER 73 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- 57024-20-1 ΙT RL: PRP (Properties)

(NMR of)

RN 57024-20-1 CAPLUS

CN 3,4-Furandimethanol, α 4,2-bis[4-(acetyloxy)phenyl]tetrahydro-, diacetate (9CI) (CA INDEX NAME)

L44 ANSWER 74 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 55665-05-9P

RL: SPN (Synthetic preparation); PREP (Preparation)
(preparation of)

RN 55665-05-9 CAPLUS

CN Methanone, [4-(acetyloxy)-3,5-dimethoxyphenyl][5-[4-(acetyloxy)-3,5-dimethoxyphenyl]-4-[(acetyloxy)methyl]tetrahydro-3-furanyl]- (CA INDEX NAME)

- L44 ANSWER 75 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 54797-70-5 54797-72-7 54849-03-5 RL: PRP (Properties)

(ir spectra of)

RN 54797-70-5 CAPLUS

CN 2(3H)-Furanone, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, (3R,4R)- (CA INDEX NAME)

Absolute stereochemistry.

54797-72-7 CAPLUS RN

CN 3,4-Furandimethanol, α,α' -bis[4-(acetyloxy)-3methoxyphenyl]tetrahydro-, diacetate (9CI) (CA INDEX NAME)

RN 54849-03-5 CAPLUS

2(3H)-Furanone, 4-[(acetyloxy)[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-[[4-CN $(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, [3S-[3<math>\alpha$, 4 β (R*)]]-(9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L44 ANSWER 76 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- 34000-77-6 RL: PRP (Properties)

IT

(carbon-13 NMR of) RN 34000-77-6 CAPLUS

CN

Phenol, 4,4'-[(tetrahydro-3,4-furandiyl)bis(methylene)]bis[2,6-dimethoxy-, diacetate (9CI) (CA INDEX NAME)

L44 ANSWER 77 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 40516-25-4P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

40516-25-4 CAPLUS RN

1,2-Benzenediol, 4-[4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-CN [(acetyloxy)methyl]tetrahydro-2-furanyl]-, diacetate (9CI) (CA INDEX NAME)

L44 ANSWER 78 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

ΙT 34000-77-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 34000-77-6 CAPLUS

CN

Phenol, 4,4'-[(tetrahydro-3,4-furandiyl)bis(methylene)]bis[2,6-dimethoxy-, diacetate (9CI) (CA INDEX NAME)

IT 30031-99-3P
 RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 30031-99-3 CAPLUS

CN 2(3H)-Furanone, 3-(3,4-dihydroxy-5-methoxybenzylidene)dihydro-4-vanillyl-, triacetate (8CI) (CA INDEX NAME)

L44 ANSWER 80 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 21497-65-4

RL: PRP (Properties)

(rearrangement (photochem.) of)

RN 21497-65-4 CAPLUS

2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (Z,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 21497-66-5

RL: PRP (Properties)

(spectrum (ir and uv) of)

RN 21497-66-5 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-, (E,Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L44 ANSWER 81 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 6512-68-1P, Thujaplicatin, O-methyl-, diacetate, trans-RL: PREP (Preparation)

(preparation of) RN 6512-68-1 CAPLUS

KN 6512-68-1 C

CN 2(3H)-Furanone, 3-[[4-(acetyloxy)-3,5-dimethoxyphenyl]methyl]-4-[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

Currently available stereo shown.

- L44 ANSWER 82 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 98770-68-4
- (Derived from data in the 6th Collective Formula Index (1957-1961))
- RN 98770-68-4 CAPLUS
- CN 2(3H)-Furanone, 4-[4-(acetyloxy)-3-methoxybenzoyl]-3-[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, (3R-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

IT 54797-72-7, 3,4-Furandimethanol, tetrahydro- α , α '-bis(4-hydroxy-3-methoxyphenyl)-, tetraacetate 909256-63-9, Hydrocinnamic acid, 4-hydroxy- α -(4-hydroxy- α -(hydroxymethyl)-3-methoxyphenacyl]-3-methoxy-, (+)-, diacetate

(from fir wood)

RN 54797-72-7 CAPLUS

CN 3,4-Furandimethanol, α,α' -bis[4-(acetyloxy)-3-methoxyphenyl]tetrahydro-, diacetate (9CI) (CA INDEX NAME)

RN 909256-63-9 CAPLUS

CN 2(3H)-Furanone, 4-[4-(acetyloxy)-3-methoxybenzoy1]-3-[[4-(acetyloxy)-3-methoxypheny1]methy1]dihydro- (CA INDEX NAME)

L44 ANSWER 83 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 98770-68-4

(Derived from data in the 6th Collective Formula Index (1957-1961))

RN 98770-68-4 CAPLUS

CN 2(3H)-Furanone, 4-[4-(acetyloxy)-3-methoxybenzoyl]-3-[[4-(acetyloxy)-3-methoxyphenyl]methyl]dihydro-, (3R-trans)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

L44 ANSWER 84 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

IT 63339-52-6P, Fulgide, 6,7-bis(4-hydroxy-3-methoxyphenyl)-,

diacetate

RL: PREP (Preparation) (preparation of)

RN 63339-52-6 CAPLUS

CN 2,5-Furandione, 3,4-bis[[4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro-(CA INDEX NAME)

L44 ANSWER 85 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN 860255-29-4P, Hydroferulic acid, α -[β , 4-dihydroxyα-(hydroxymethyl)-3-methoxyphenethyl]-β-hydroxy-, γ-lactone, tetraacetate

RL: PREP (Preparation)

(preparation of) 860255-29-4 CAPLUS

RN CN

2(3H)-Furanone, 4-[(acetyloxy)[4-(acetyloxy)-3-methoxyphenyl]methyl]-3-[(acetyloxy) [4-(acetyloxy)-3-methoxyphenyl]methylene]dihydro- (CA INDEX NAME)

L44 ANSWER 86 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN

910882-90-5P, Matairesinol, bis(p-nitrobenzoate)

RL: PREP (Preparation) (preparation of)

RN 910882-90-5 CAPLUS

2(3H)-Furanone, dihydro-3,4-bis[[3-methoxy-4-[(4nitrobenzoyl)oxy]phenyl]methyl]-, (3R,4R)- (CA INDEX NAME)

Absolute stereochemistry.

PAGE 1-A

PAGE 1-B

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L36

They are available for your review at: http://www.cas.org/legal/infopolicy.html => d his (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008) FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON 1 US20050249857/PN FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 L2 TRA L1 1- RN : 16 TERMS FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 L3 16 SEA L2 L4 10 L3 AND >=2 46.150.18/RID 1 L3 AND OC4-C6-C6/ES L5 L6 1 L3 AND C6-C6/ES L7 STR 1.8 29 L7 L9 SCR 1839 29 L7 AND L9 L10 L11 STR L7 L12 336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES L13 50 L11 SAM SUB=L12 L14 STR L11 32 L14 SAM SUB=L12 L15 E FURAN/CN 1 E3 L16 E THF/CN L17 1 E3 L18 130953 L12 AND 16.138.1/RID L19 49 L14 SAM SUB=L18 L20 STR L14 FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008 FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008 L21 STR L20 L22 SCR 1707 OR 1708 L23 SCR 1840 L24 2 L22 AND L23 AND L21 L25 36 L22 AND L23 AND L21 SAM SUB=L18 L26 STR L21 L27 4 L26 AND L22 AND L23 SAM SUB=L18 L28 94 L26 AND L22 AND L23 FULL SUB=L18 SAV TEM G761C1/A L28 L29 STR L7 L30 2 L29 L31 STR L29 L32 0 L31 SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705 1.33 L34 0 L31 AND L33 L35 33 L31 AND L33 FULL

SAV TEM G761C1N/A L35

0 L28, L35 AND L3

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L38
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L41
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     FILE 'CAPLUS' ENTERED AT 15:30:02 ON 12 JUN 2008
=> d 144 60
L44 ANSWER 60 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
     1980:8015 CAPLUS
AN
DN
     92:8015
OREF 92:1465a,1468a
    NMR studies of lignins. 3. Proton NMR spectroscopic data for lignin
     model compounds
     Lundquist, Knut
AU
CS
     Dep. Org. Chem., Chalmers Univ. Technol., Goeteborg, S-412 96, Swed.
SO
     Acta Chemica Scandinavica, Series B: Organic Chemistry and Biochemistry
     (1979), B33(6), 418-20
     CODEN: ACBOCV; ISSN: 0302-4369
DT
     Journal
LA
     English
=> d bib abs hitstr 60
L44 ANSWER 60 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
AN
    1980:8015 CAPLUS
DN
    92:8015
OREF 92:1465a,1468a
    NMR studies of lignins. 3. Proton NMR spectroscopic data for lignin
TΤ
     model compounds
AII
     Lundquist, Knut
CS
     Dep. Org. Chem., Chalmers Univ. Technol., Goeteborg, S-412 96, Swed.
     Acta Chemica Scandinavica, Series B: Organic Chemistry and Biochemistry
SO
     (1979), B33(6), 418-20
     CODEN: ACBOCV; ISSN: 0302-4369
DT
     Journal
LA
     English
AB
     The 1H NMR spectra of a number of lignin (I) [9005-53-2] model compds.
     showed that the \alpha-H signal is shifted by 0.03-0.06 \delta units
     upfield when an adjacent 4-acetoxy-3-methoxyphenyl group in the model
     compound is replaced by a 3,4-dimethoxyphenyl (II) group (representative of
     4-alkoxy-3-methoxyphenyl groups in I). The \alpha-H in
     4-acetoxy-3-methoxybenzyl acetate [17574-14-0] and 6,6'-dihydroxy-5,5'-
     dimethoxy-[1,1'-bipheny1]-3,3'-dimethanol tetraacetate [72092-47-8]
     exhibited about the same \delta value, indicating that the introduction
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of a biphenyl linkage does not change the position of the $\alpha\text{-H}$ signal to any great extent. erythro And three forms of model compds. differed moderately in their 1H NMR properties.

IT 72092-51-4

RL: PRP (Properties)

(NMR spectrum of, as model for lignin)

RN 72092-51-4 CAPLUS

CN Phenol, 4,4'-[(tetrahydro-3,4-furandiyl)bis(methylene)]bis[2-methoxy-, diacetate, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

=> FIL STNGUIDE

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- L44 ANSWER 2 OF 86 CAPLUS COPYRIGHT 2008 ACS on STN
- AN 2007:17912 CAPLUS
- DN 147:318338
- TI Structure of lignins in developing xylem of Norway spruce
- AU Christiernin, M.
- CS Department of Fiber and Polymer Technology, Royal Institute of Technology, KTH, Stockholm, 100 44, Swed.
- SO Plant Physiology and Biochemistry (Amsterdam, Netherlands) (2006), 44(11-12), 693-699 CODEN: PPBIEX: ISSN: 0981-9428
 - CODEN: PPBIEX;
- PB Elsevier B.V.
- DT Journal
- LA English
 AB The developing xylem in a Norway spruce (Picea abies) clone was
- investigated during a growth season and compared to lignin from sapwood of the same tree clone. Klason and acid-soluble lignin contents were determined

well as the carbohydrate monomer distribution and protein content. By analyzing lignin thioacidolysis products, it was shown that only gualacel units could be detected in the materials, and the relative amount of $\beta\text{-}O\text{-}4\text{'}$ bonds was assessed. Monomeric and selected dimeric lignin products were identified by mass spectrometry. The specimens were embedded and thin sections examined by microscopy to determine the state of

cell

differentiation in the samples. In the spring and early summer, growth was very rapid and the intention was to collect tissue in which exclusively the middle lamella/primary cell wall had begun to lignify. Combining data regarding Klason lignin, protein content and carbohydrate monomer distribution with microscopy, it was found that the developing xylem sample from mid-June contained lignin from exclusively middle lamella/primary wall. The Klason lignin content in the developing xylem during the growth season was 20%, 5% and 10% in Apr., June and August, resp. Thioacidolysis showed that the lignin had more condensed structures than lignin from the reference Norway spruce clone wood. Mass spectrometry showed that the developing xylem specimens from June and August contained more lignin structures with end-groups than the reference sample. These results suggest that lignification in the cambial layer and early developing xylem may take place more in a bulk fashion during the summer.

IT 947685-66-7

RL: BSU (Biological study, unclassified); BIOL (Biological study) (content of lignin; structure of lignins in developing xylem of Norway spruce (Picea abies))

RN 947685-66-7 CAPLUS

CN Phenol, 4,4'-[(tetrahydro-3,4-furandiyl)bis(methylene)]bis[2-methoxy-, 1,1'-diacetate (CA INDEX NAME)

RE.CNT 26 THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d bib abs hitstr 7
YOU HAVE REQUESTED DATA FROM FILE 'CAPLUS' - CONTINUE? (Y)/N:y

- DN 128:153961
- TI Interaction of lignans with human sex hormone-binding globulin (SHBG)
- AU Schoettner, Matthias; Gansser, Dietmar; Spiteller, Gerhard
- CS Universitaet Bayreuth, Bayreuth, D-95440, Germany
- SO Zeitschrift fuer Naturforschung, C: Biosciences (1997), 52(11/12), 834-843 CODEN: ZNCBDA; ISSN: 0341-0382
 - Verlag der Zeitschrift fuer Naturforschung
- PB Verlag de DT Journal
- LA English
- AB In a double Stobbe condensation without use of protecting groups a wide variety of lignans with different substitution pattern in the aromatic and aliphatic part of the mol. was synthesized. These lignans were tested in a sex normone-binding globulin binding assay which allowed to deduce the following relationship between structure and activity:
 - (±)-diastereomers are more active than meso compds., the 4-hydroxy 3-methoxy (quajacyl) substitution pattern in the aromatic part is most effective, and the activity increases with the decline in polarity of the aliphatic part of the mol.
- IT 72092-51-4
 - RL: BSU (Biological study, unclassified); PEP (Physical, engineering or chemical process); BIOL (Biological study); PROC (Process) (preparation of lionans and interaction with sex hormone-binding globulin)
- RN 72092-51-4 CAPLUS
- CN Phenol, 4,4'-[(tetrahydro-3,4-furandiy1)bis(methylene)]bis[2-methoxy-, diacetate, trans- (9CI) (CA INDEX NAME)

Relative stereochemistry.

=> d his

(FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008)

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1.1 1 IIS20050249857/PN

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- L4 10 L3 AND >=2 46.150.18/RID
- L5 1 L3 AND OC4-C6-C6/ES
- L6 1 L3 AND C6-C6/ES
- L7 STR
- L8 29 L7
- L9 SCR 1839
- L10 29 L7 AND L9
- L11 STR L7

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L14
                STR L11
L15
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L16
              1 E3
               E THF/CN
L17
              1 E3
L18
        130953 L12 AND 16.138.1/RID
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L19
L20
                STR L14
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L22
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L23
                SCR 1840
L24
              2 L22 AND L23 AND L21
L25
             36 L22 AND L23 AND L21 SAM SUB=L18
L26
                STR L21
L27
              4 L26 AND L22 AND L23 SAM SUB=L18
L28
             94 L26 AND L22 AND L23 FULL SUB=L18
                SAV TEM G761C1/A L28
1.29
                STR L7
L30
              2 L29
L31
                STR L29
L32
             0 L31
                SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705
L33
L34
              0 L31 AND L33
L35
             33 L31 AND L33 FULL
                SAV TEM G761C1N/A L35
L36
              0 L28, L35 AND L3
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     FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
L37
                STR
L38
             46 L37
L39
        694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES
L40
             50 L37 SAM SUB=L39
L41
                STR L37
L42
             50 L41 SAM SUB=L39
L43
           7953 L41 FULL SUB=L39
                SAV TEM G761C1N2/A L43
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L44
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     FILE 'CAPLUS' ENTERED AT 15:30:02 ON 12 JUN 2008
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FILE 'STNGUIDE' ENTERED AT 15:37:57 ON 12 JUN 2008

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=> s 135 L45 55 T-35

=> d hitstr 1-55

- L45 ANSWER 1 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IΤ 892403-74-6P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of vitamin E intermediates, phytyl hydroquinone derivs. by cross-metathesis of allylhydroquinones with tetramethylhexadecenyl esters and aldehyde)

892403-74-6 CAPLUS RN

1,4-Benzenediol, 2,2'-(2-butene-1,4-diyl)bis(3,5,6-trimethyl-, CN 4,4'-diacetate (CA INDEX NAME)

- L45 ANSWER 2 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 892403-74-6P

RL: BYP (Byproduct); PREP (Preparation)

(preparation of olefins as vitamin E precursors by cross-metathesis) RN 892403-74-6 CAPLUS

CN 1,4-Benzenediol, 2,2'-(2-butene-1,4-divl)bis[3,5,6-trimethyl-, 4,4'-diacetate (CA INDEX NAME)

- L45 ANSWER 3 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 180002-86-2

RL: MOA (Modifier or additive use); USES (Uses)

(compns. of stabilizers in resins for battery or capacitor separators)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

- L45 ANSWER 4 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 823808-17-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis and thermal properties of main chain polyimides containing chalcone derivative)

- RN 823808-17-9 CAPLUS
- CN Benzoic acid, 4-[(11-hydroxyundecyl)oxy]-, 4-[3-[4-[(1-[(1-hydroxyundecyl)oxy]phenyl]-1-oxo-2-propen-1-yl]phenyl ester (CA INDEX NAME)

PAGE 1-B

- L45 ANSWER 5 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 180002-86-2, GSY 242

RL: MOA (Modifier or additive use); USES (Uses)

(antioxidant; shaped articles of antioxidant-containing polyolefin resin compns. and vinyl chloride resins)

- RN 180002-86-2 CAPLUS
- CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-o-benylene]] ester (CA INDEX NAKE)

t-Bu

PAGE 2-B

- L45 ANSWER 6 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 180002-86-2
 - RL: MOA (Modifier or additive use); USES (Uses)
 - (stabilized polymer composition containing hindered phenols, phosphorus and sulfur compds.)
- RN 180002-86-2 CAPLUS
- CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-ohenylenel] ester (CA INDEX NAME)

PAGE 1-A

t-Bu

L45 ANSWER 7 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

T 180002-86-2 180002-87-3 202331-18-8

202331-19-9 RL: MOA (Modifier or additive use); USES (Uses)

(stabilized polyolefin resin composition containing low concentration of antioxidant)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

PAGE 1-A

t-Bu

PAGE 2-A

PAGE 2-B

RN 180002-87-3 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1-phenylene] ester (9C1) (CA INDEX NAME)

t-Bu

PAGE 2-B

t-Bu

CN Benzenepropanoic acid, 3-(1,1-dimethylethyl)-4-hydroxy-5-methyl-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

RN 202331-19-9 CAPLUS

NN 22331-19-7 CAFEDO acid, 3-(1,1-dimethylethyl)-4-hydroxy-5-methyl-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1-phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

t-Bu

PAGE 2-A

PAGE 2-B

L45 ANSWER 8 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 339588-47-5P

RL: SPN (Synthetic preparation); PREP (Preparation) (solid-phase synthesis of alkadiynediol bis(benzofurancarboxylate)s via acetylenic homocoupling catalyzed by AgOTS-CuC12-TMEDA)

RN 339588-47-5 CAPLUS

CN Benzenemethanol, 3,3'-(1,3-butadiyne-1,4-diyl)bis[4-(acetyloxy)-5-methoxy-(CA INDEX NAME)

- L45 ANSWER 9 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
 - T 180002-86-2 180002-87-3 202331-18-8

202331-19-9

RL: NOA (Modifier or additive use); USES (Uses) (degradation inhibitor for resin material, chlorine water-resistant resin composition and method for inhibiting degradation)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-

4,1-phenylene]] ester (CA INDEX NAME)

PAGE 1-A

t-Bu

PAGE 2-B

RN 180002-87-3 CAPLUS

Name of the state of the state

PAGE 1-A

RN 202331-18-8 CAPLUS

NN 20233-10-0 CAEDOG Acid, 3-(1,1-dimethylethyl)-4-hydroxy-5-methyl-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

$$\begin{array}{c} \text{HO} \\ \text{Me} \\ \text{Me} \end{array} \\ \begin{array}{c} \text{CH}_2 \\ \text{CH}_2 \\ \text{CH} \\ \text{CH}_2 \\ \text{CH} \\ \text{CH}_2 \\ \text{$$

PAGE 2-B

RN 202331-19-9 CAPLUS

CN Benzenepropanoic acid, 3-(1,1-dimethylethyl)-4-hydroxy-5-methyl-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1-phenylene] ester (9CI) (CA INDEX NAME)

PAGE 2-B

L45 ANSWER 10 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 286382-99-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(effect of phosphorylated phloretin derivative effect intestinal Na+-dependent phosphate absorption)

RN 286382-99-8 CAPLUS

CN 1-Propanone, 3-[4-(acetyloxy)phenyl]-1-[2,4-bis(acetyloxy)-6-(β-D-glucopyranosyloxy)phenyl]- (CA INDEX NAME)

Absolute stereochemistry.

L45 ANSWER 11 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 180002-86-2, GSY 242

RL: MOA (Modifier or additive use); USES (Uses)

(GSY 242; environmental change surrounding resin antioxidants)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

L45 ANSWER 12 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 401796-81-4P 401796-82-5P

RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)

(pigment; preparation of carthamin acetate red pigment in safflower petals) RN 401796-81-4 CAPLUS

CN 4-Cyclohexene-1,3-dione, 6-(acetyloxy)-2-[[(3S)-3-(acetyloxy)-5-[(2E)-3-[4-(acetyloxy)]henyl]-1-oxo-2-propenyl]-2,4-dihydroxy-6-oxo-3-((2R,3S,4R,5S,6S)-3,4,5-tris(acetyloxy)-6-[(acetyloxy)methyl]tetrahydro-2H-pyran-2-yl]-1,4-cyclohexadien-1-yl]methylene]-4-[(2E)-3-[4-(acetyloxy)phenyl]-1-oxo-2-propenyl]-5-hydroxy-6-[(2E,3R,4S,5R,6R)-3,4,5-tris(acetyloxy)-6-[(acetyloxy)methyl]tetrahydro-2H-pyran-2-yl]-, (2Z,6S)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

RN 401796-82-5 CAPLUS

CN 4-Cyclohexene-1,3-dione, 6-(acetyloxy)-2-[[(3R)-3-(acetyloxy))-5-[(2E)-3-[4-(acetyloxy))hemyl]-1-cox-2-propenyl]-2,4-dihydroxy-6-cox-3-[(2R,3S,4R,5S,6S)-3,4,5-tris(acetyloxy)-6-[(acetyloxy)methyl]tetrahydro-2Hpyran-2-yl]-1,4-cyclohexadien-1-yl]methylene]-4-[(2E)-3-[4-(acetyloxy)hemyl]-1-cox-2-propenyl]-5-hydroxy-6-[(2S,3R,45,5R,6R)-3,4,5tris(acetyloxy)-6-((acetyloxy)methyl]tetrahydro-2H-pyran-2-yl]-, (2Z,6R)-(9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L45 ANSWER 13 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

339588-47-5P ΙT RL: BYP (Byproduct); SPN (Synthetic preparation); PREP (Preparation) (optimization of Sonogashira cross-coupling on high-loading macrobeads using silvl linker)

339588-47-5 CAPLUS CN

Benzenemethanol, 3,3'-(1,3-butadiyne-1,4-diyl)bis[4-(acetyloxy)-5-methoxy-(CA INDEX NAME)

- L45 ANSWER 14 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- 180002-86-2 ΙT

RN

RL: DEV (Device component use); MOA (Modifier or additive use); USES (Uses)

(thermal printing material containing phenolic compound decoloration preventing agent)

180002-86-2 CAPLUS RN

Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, CN 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

L45 ANSWER 15 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 286382-99-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction; aryl phosphate, thiophosphate, and aminophosphate inhibitors of intestinal apical membrane sodium/phosphate co-transport, and therapeutic use)

RN 286382-99-8 CAPLUS

CN 1-Propanone, 3-[4-(acetyloxy)phenyl]-1-[2,4-bis(acetyloxy)-6-(β-D-glucopyranosyloxy)phenyl]- (CA INDEX NAME)

Absolute stereochemistry.

L45 ANSWER 16 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 252353-00-7P

RL: DEV (Device component use); NUU (Other use, unclassified); PNU (Preparation, unclassified); PREP (Preparation); USES (Uses) (preparation of optically active butane-tetrol derivative for doping of

liquid

crystals in display devices) RN 252353-00-7 CAPLUS

CN Benzoic acid, 4-[[6-[(1-oxo-2-propenyl)oxy]hexyl]oxy]-, (2R,3R)-1,2,3,4-tetrahydroxy-1,4-butanediylidene]tetra-4,1-phenylene ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

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PAGE 1-B

PAGE 2-B

L45 ANSWER 17 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 180002-86-2 180002-87-3 202331-18-8 242467-79-4

RL: MOA (Modifier or additive use); USES (Uses)

(antioxidant; flame retardant polyolefin compns. with good heavy metal-induced oxidation resistance)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(n-ethyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

PAGE 1-A

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PAGE 2-B

RN 180002-87-3 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propan)-3-ylidene)tris(2-(1,1-dimethylethyl)-6-methyl-4,1-phenylene) ester (9CI) (CA INDEX NAME)

t-Bu

PAGE 2-B

t-Bu

CN Benzenepropanoic acid, 3-(1,1-dimethylethyl)-4-hydroxy-5-methyl-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

RN 242467-79-4 CAPLUS

NN 24240 - 7-7-4 CAEDOO COME SET OF THE SET

PAGE 1-A

Me | Me-C-Et

PAGE 2-B

PAGE 2-A HO Me CH-Me CH-Me CH-Me
$$CH_2$$
 CH CH_2 CH C

Мe C-Et Me

- L45 ANSWER 18 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN IT $180002{-}86{-}2$ $180002{-}87{-}3$
- - RL: MOA (Modifier or additive use); PRP (Properties); USES (Uses) (antioxidant; polyolefin pipes with excellent oxidative degradation resistance in the presence of water)
- 180002-86-2 CAPLUS RN

PAGE 1-A

PAGE 2-A

CN

RN 180002-87-3 CAPLUS

Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1phenylene] ester (9CI) (CA INDEX NAME)

PAGE 1-A

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PAGE 2-B

- L45 ANSWER 19 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 180002-86-2 180002-87-3 202331-18-8 242467-79-4

RL: MOA (Modifier or additive use); USES (Uses)
(stabilized polyolefin compns. for use in heat-exchange system)

- RN 180002-86-2 CAPLUS
- CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c} \text{HO} \\ \text{HO} \\ \text{CH}_2 - \text{CH}_2 - \text{CH}_2 - \text{C} \\ \text{O} \\ \text{Me} \\ \text{Me} \\ \text{CH} \\ \text{CH} \\ \text{O} - \text{C} - \text{CH}_2 - \text{CH}_2 \\ \text{CH}_2 \\ \text{O} \\ \text{C} + \text{Du} \\ \text{O} \\ \text{C} + \text{Du}$$

RN 202331-18-8 CAPLUS

NN 20233-10-0 CAEDOG Acid, 3-(1,1-dimethylethyl)-4-hydroxy-5-methyl-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

HO O T-Bu CH-Me CH₂ CH₂ Me CH₂ Me
$$CH_2$$
 Me CH_2 Me CH_2 Me CH_2 Me CH_2 Me CH_2 Me

PAGE 2-B

RN 242467-79-4 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylpropyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2,6-bis(1,1-dimethylethyl)-4,1phenylene] ester (9C1) (CA INDEX NAME)

Me-C-Et

PAGE 1-B

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PAGE 2-B

L45 ANSWER 20 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 180002-86-2 180002-87-3

RL: MOA (Modifier or additive use); USES (Uses) (antioxidants; durable crosslinked polyolefin tubes containing antioxidants including hindered phenols)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

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L45 ANSWER 21 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 180002-86-2P

RL: IMF (Industrial manufacture); MOA (Modifier or additive use); PREP (Preparation); USES (Uses)

(antioxidants; water-resistant extrusion moldings of crosslinked polyolefin resins and manufacture)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-,
1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl4,1-phenylene]] ester (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

PAGE 2-B

L45 ANSWER 22 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 180002-86-2P

RL: IMF (Industrial manufacture); MOA (Modifier or additive use); PREP (Preparation); USES (Uses) (polyolefin composition with good mech. strength and good discoloration

prevention after in hot or cool water for a long period) RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

L45 ANSWER 23 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

180002-86-2P

RL: IMF (Industrial manufacture); MOA (Modifier or additive use); PREP (Preparation); USES (Uses)

(manufacture of beta-crystalline oligomeric hindered phenolic compound for use as

antioxidant)

180002-86-2 CAPLUS RN

Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, CN

1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-

4,1-phenylene]] ester (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c} \text{HO} & \text{O} & \text{CH} - \text{Me} \\ \text{CH}_2 - \text{CH}_2 - \text{C} - \text{O} & \text{CH} - \text{Me} \\ \text{CH}_2 & \text{CH}_2 - \text{CH}_2$$

PAGE 2-B

L45 ANSWER 24 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

156728-76-6P 156728-77-7P 156728-78-8P тт

156728-79-9P 156728-80-2P RL: BAC (Biological activity or effector, except adverse); BSU (Biological

study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of dihydrochalcone sugar derivs. as antidiabetic agents)

CN

156728-76-6 CAPLUS
1-Propanone, 1-[2-(acetyloxy)-6-(β-D-glucopyranosyloxy)phenyl]-3-[4-(acetyloxy)phenyl]- (CA INDEX NAME)

Absolute stereochemistry.

156728-77-7 CAPLUS RN

CN Propanoic acid, 2-methyl-, 3-(β-D-glucopyranosyloxy)-2-[3-[4-(2methyl-1-oxopropoxy)phenyl]-1-oxopropyl]phenyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 156728-78-8 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, 3-(β-D-glucopyranosyloxy)-2-[3-[4-[[(2-methylpropoxy)carbonyl]oxy]phenyl]-1-oxopropyl]phenyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 156728-79-9 CAPLUS

CN Carbonic acid, 4-[3-[2-[(ethoxycarbonyl)oxy]-6-(β-D-glucopyranosyloxy)phenyl]-3-oxopropyl]phenyl ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 156728-80-2 CAPLUS

CN Carbonic acid, 3-(β-D-glucopyranosyloxy)-2-[3-[4-[[(2-methylpropoxy)carbonyl]oxy]phenyl]-1-oxopropyl]phenyl 2-methylpropyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- L45 ANSWER 25 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 180002-86-2 180002-87-3

RL: MOA (Modifier or additive use); PRP (Properties); TEM (Technical or engineered material use); USES (Uses)

(bleeding-resistant durable polyolefin injection moldings containing antioxidants for jacketting steel pipes)

- RN 180002-86-2 CAPLUS
- CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

t-Bu

L45 ANSWER 26 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

T 180002-86-2 180002-87-3 202331-18-8 202331-19-9

RL: MOA (Modifier or additive use); TEM (Technical or engineered material use); USES (Uses)

(polyolefin compns. with good hot water resistance)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-,
1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl4,1-phenylene]] ester (CA INDEX NAME)

PAGE 1-A

$$\begin{array}{c} \text{OH} \\ \text{Eu-t} \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{C} \\ \text{C} \end{array}$$

t-Bu

PAGE 2-A

PAGE 2-B

RN 180002-87-3 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1-phenylene] ester (9C1) (CA INDEX NAME)

t-Bu

PAGE 2-B

t-Bu

CN Benzenepropanoic acid, 3-(1,1-dimethylethyl)-4-hydroxy-5-methyl-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

RN 202331-19-9 CAPLUS

NN 22331-19-7 CAFLOOR CAPUAGE AND A NAMED AND A NAMED CAPTURE CAPTURE

PAGE 1-A

OH

t-Bu

PAGE 2-A

$$\begin{array}{c} \text{HO} \\ \text{Me} \end{array} \begin{array}{c} \text{CH}_{-\text{Bu}} \\ \text{CH}_{2} \\ \text{CH} \end{array} \begin{array}{c} \text{CH}_{-\text{Me}} \\ \text{CH}_{2} \\ \text{CH} \\ \text{O}_{-\text{C}} \\ \text{CH}_{2} \\ \text{CH}_{2} \end{array} \begin{array}{c} \text{Me} \\ \text{O}_{-\text{C}} \\ \text{CH}_{2} \\ \text{CH}_{2} \\ \text{O}_{-\text{C}} \end{array}$$

PAGE 2-B

L45 ANSWER 27 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 180002-86-2

RL: MOA (Modifier or additive use); USES (Uses)
(antioxidants; polypropylene compns. containing hindered phenols with good durability for steel pipe coatings)

RN 180002-86-2 CAPLUS

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

- L45 ANSWER 28 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 180002-86-2 180002-87-3

RL: MOA (Modifier or additive use); USES (Uses) (polyolefine containing hindered phenols as antioxidants for extrusion moldings)

- RN 180002-86-2 CAPLUS
- CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

PAGE 1-A

t-Bu

CN

RN 180002-87-3 CAPLUS

Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1-phenylene] ester (9CI) (CA INDEX NAME)

PAGE 1-A

t-Bu

PAGE 2-A

$$\begin{array}{c} \text{HO} \\ \text{t-Bu} \\ \text{t-Bu} \\ \end{array} \begin{array}{c} \text{CH-Me} \\ \text{CH}_2 \\ \text{C$$

PAGE 2-B

- L45 ANSWER 29 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 182928-76-3P 182928-78-5P
- RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 - (synthesis of (+)- and (-)-model compds. and absolute configuration of carthamin)
- RN 182928-76-3 CAPLUS
- CN 2,5-Cyclohexadien-1-one, 4-(acetyloxy)-2-[3-[4-(acetyloxy)phenyl]-1-oxo-2-propenyl]-3,5-dihydroxy-4-methyl-, [R-(E)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

- RN 182928-78-5 CAPLUS
- CN 2,5-Cyclohexadien-1-one, 4-(acetyloxy)-2-[3-[4-(acetyloxy)pheny1]-1-oxo-2-propeny1]-3,5-dihydroxy-4-methyl-, [S-(E)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L45 ANSWER 30 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

174216-56-9

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study) (inhibition of cytokine production in human and rat macrophages by

dicatechol rooperol and its esters) RN 174216-56-9 CAPLUS

CN 1,2-Benzenediol, 4,4'-(1-penten-4-vne-1,5-divl)bis-, 1,1'-diacetate, (E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

- L45 ANSWER 31 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 180002-86-2P 180002-87-3P

RL: SPN (Synthetic preparation); TEM (Technical or engineered material use); PREP (Preparation); USES (Uses)

(preparation of phenolic compds. as antioxidants) RN 180002-86-2 CAPLUS

CN

Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, 1,1'-[(1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-5-methyl-4,1-phenylene]] ester (CA INDEX NAME)

t-Bu

PAGE 2-B

CN Benzenepropanoic acid, 3,5-bis(1,1-dimethylethyl)-4-hydroxy-, (1-methyl-1-propanyl-3-ylidene)tris[2-(1,1-dimethylethyl)-6-methyl-4,1phenylene] ester (9C1) (CA INDEX NAME)

PAGE 1-A

PAGE 2-A

t-Bu

PAGE 2-B

L45 ANSWER 32 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 174216-56-9P

RN

RL: BAC (Biological activity or effector, except adverse); BOC (Biological occurrence); BSU (Biological study, unclassified); PUR (Purification or recovery); THU (Therapeutic use); BIOL (Biological study); OCCU (Occurrence); PREP (Preparation); USES (Uses)

(rooperol and hypoxoside and their derivs. in treatment of inflammation)

174216-56-9 CAPLUS

CN 1,2-Benzenediol, 4,4'-(1-penten-4-yne-1,5-diyl)bis-, 1,1'-diacetate, (E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

L45 ANSWER 33 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 156728-76-6P 156728-77-7P 156728-78-8P

156728-79-9P 156728-80-2P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of glucopyranosyldihydrochalcone derivs. as hypoglycemic agents)

RN 156728-76-6 CAPLUS

CN 1-Propanone, 1-[2-(acetyloxy)-6-(β-D-glucopyranosyloxy)phenyl]-3-[4-(acetyloxy)phenyl]- (CA INDEX NAME)

RN 156728-77-7 CAPLUS

CN Propanoic acid, 2-methyl-, 3-(B-D-glucopyranosyloxy)-2-[3-[4-(2-methyl-1-oxopropoxy)phenyl]-1-oxopropyl]phenyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 156728-78-8 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, 3-(β-D-glucopyranosyloxy)-2-[3-[4-[[(2-methylpropoxy)carbonyl]oxy]phenyl]-1-oxopropyl]phenyl ester (CA INDEX NAME)

RN 156728-79-9 CAPLUS

CN Carbonic acid, 4-[3-[2-[(ethoxycarbonyl)oxy]-6-(β-D-glucopyranosyloxy)phenyl]-3-oxopropyl]phenyl ethyl ester (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 156728-80-2 CAPLUS

CN Carbonic acid, 3-(\$\Gamma-0\)-glucopyranosyloxy)-2-[3-[4-[[(2-methylpropoxy]carbony]]oxy]phenyl]-1-oxopropyl]phenyl 2-methylpropyl ester (\$\Gamma(1)\) (CA INDEX NAME)

L45 ANSWER 34 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

156728-76-6P 156728-77-7P 156728-78-8P 156728-79-9P 156728-80-2P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as hypoglycemic)

RN

156728-76-6 CAPLUS 1-Propanone, $1-[2-(acetyloxy)-6-(\beta-D-glucopyranosyloxy)phenyl]-3-[4-(b-D-glucopyranosyloxy)phenyloxy]-3-[4-(b-D-glucopyranosyloxy)phenyloxy]-3-[4-(b-D-glucopyranosyloxy)phenyloxy]-3-[4-(b-D-glucopyranosyloxy)phenyloxy]-3-[4-($ CN (acetyloxy)phenyl]- (CA INDEX NAME)

Absolute stereochemistry.

RN 156728-77-7 CAPLUS

Propanoic acid, 2-methyl-, 3-(β-D-glucopyranosyloxy)-2-[3-[4-(2-CN methyl-1-oxopropoxy)phenyl]-1-oxopropyl]phenyl ester (CA INDEX NAME)

RN 156728-78-8 CAPLUS

CN Propanoic acid, 2,2-dimethyl-, 3-(β-D-glucopyranosyloxy)-2-[3-[4-[[(2-methylpropoxy)carbonyl]oxy]phenyl]-1-oxopropyl]phenyl ester (CA INDEX NAME)

Absolute stereochemistry.

RN 156728-79-9 CAPLUS

CN Carbonic acid, 4-[3-[2-[(ethoxycarbonyl)oxy]-6-(β-D-glucopyranosyloxy)phenyl]-3-oxopropyl]phenyl ethyl ester (9CI) (CA INDEX NAME)

RN 156728-80-2 CAPLUS

CN Carbonic acid, 3-(β-D-glucopyranosyloxy)-2-[3-[4-[[(2-methylpropoxylcarbonyl]oxy]phenyl]-1-oxopropyl]phenyl 2-methylpropyl ester (9CI) (CA INDEX NAME)

- L45 ANSWER 35 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 149475-44-5P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)
- RN 149475-44-5 CAPLUS
- CN 3,4-Pyrrolidinediol, 1-acetyl-2-[[6-(acetyloxy)-4-[3-[4-(acetyloxy)phenyl]1-oxo-2-propenyl]-5-hydroxy-3-oxo-6-(2,3,4,6-tetra-0-acetyl-B-Dglucopyranosyl)-1,4-cyclohexadien-1-yl]hydroxymethylene]-5[(acetyloxy)methyl]-, 3,4-diacetate (9CI) (CA INDEX NAME)

L45 ANSWER 36 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

ΙT 128373-97-7

RL: PRP (Properties) (cytotoxicity of, in L1210 cells)

128373-97-7 CAPLUS RN

CN 1,3,5-Benzenetriol, 2-[3-[3-(acetyloxy)-4-methoxyphenyl]propyl]-, 5-acetate (CA INDEX NAME)

L45 ANSWER 37 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT

RL: FORM (Formation, nonpreparative); PREP (Preparation) (formation of, by reduction of flavanone with cyanoborohydride in trifluoroacetic acid)

RN 128373-97-7 CAPLUS CN

1,3,5-Benzenetriol, 2-[3-[3-(acetyloxy)-4-methoxyphenyl]propyl]-, 5-acetate (CA INDEX NAME)

L45 ANSWER 38 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN ΙT 77782-94-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

77782-94-6 CAPLUS RN

CN 4H-1-Benzopyran-4-one, 2-[4-[2-(acetyloxy)-5-[3-[4-(acetyloxy)phenyl]-1 $oxo-2-propenyl]-4,6-dihydroxy-3-(2,3,4,6-tetra-0-acetyl-\beta-D$ glucopyranosyl)phenoxy[phenyl]-2,3-dihydro-5,7-dihydroxy-, [S-(E)]- (9CI) (CA INDEX NAME)

L45 ANSWER 39 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 62605-86-1

> RL: MOA (Modifier or additive use); USES (Uses) (heat stabilizers, containing thiodipropionic acid esters, for polymers)

RN 62605-86-1 CAPLUS

Carbonic acid, 4-[3,3-bis[4-[[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-1-methylpropyl]-2-(1,1dimethylethyl)-5-methylphenyl 4-[3-[4-[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5methylphenoxy[carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-1-[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenyl ester (9CI) (CA INDEX NAME)

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L45 ANSWER 40 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 81436-91-1P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 81436-91-1 CAPLUS

CN 1,4-Butanedio1, 2,3-bis[[3-(benzoyloxy)phenyl]methyl]-, (R*,R*)- (9CI)
 (CA INDEX NAME)

Relative stereochemistry.

- L45 ANSWER 41 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 79055-13-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent) (preparation and hydrolysis of)

- RN 79055-13-3 CAPLUS
- CN 2-Propenoic acid, 3-(4-hydroxy-3-methoxyphenyl)-, 2,3-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]-1,4-butanediyl ester, [R-(R*,R*)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry unknown.

- L45 ANSWER 42 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 77782-94-6P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

- RN 77782-94-6 CAPLUS
- CN 4H-1-Benzopyran-4-one, 2-[4-[2-(acetyloxy)-5-[3-[4-(acetyloxy)phenyl]-1-oxo-2-propenyl]-4,6-dihydroxy-3-(2,3,4,6-tetra-0-acetyl-β-D-glucopyranosyl)phenoxylphenyl]-2,3-dihydro-5,7-dihydroxy-, [S-(E)]- (9CI) (CA INDEX NAME)

- L45 ANSWER 43 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN ΙT 62605-86-1
 - RL: USES (Uses)

 - (heat stabilizers containing, for thermoplastics)
- 62605-86-1 CAPLUS RN
- CN Carbonic acid, 4-[3,3-bis[4-[[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-1-methylpropyl]-2-(1,1dimethylethyl) -5-methylphenyl 4-[3-[4-[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5methylphenoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-1-[5-
 - (1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenyl ester (9CI) (CA INDEX NAME)

PAGE 2-B

PAGE 2-C

- L45 ANSWER 44 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 62605-86-1 RL: MOA (Modifier or additive use); USES (Uses)
 - (heat stabilizers, containing thiodipropionic acid polyesters, for polymers)
- RN 62605-86-1 CAPLUS

 $\label{eq:dimethylethyl} $$ \dim thyle thyl phenyl $4-3-[4-[[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methyl phenyl] butyl]-2-(1,1-dimethylethyl)-5- methyl phenoxy [carbonyl] oxyl-5-(1,1-dimethylethyl)-2-methyl phenyl]-1-[5-(1,1-dimethylethyl)-4-hydroxy-2-methyl phenyl] butyl]-2-(1,1-dimethylethyl)-5-methyl phenyl ester (9CI) (CA INDEX NAME)$

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- L45 ANSWER 45 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 62605-86-1

RL: USES (Uses)

(heat and light stabilizers, with piperidine derivs., for thermoplastic resins)

RN 62605-86-1 CAPLUS

 $\begin{array}{ll} \mathbb{CN} & \mathsf{Carbonic} \ \mathsf{acid}, \ 4-[3,3-\mathsf{bis}[4-[[[4-[1,3-\mathsf{bis}[5-(1,1-\mathsf{dimethylethyl})-4-\mathsf{hydroxy-2-methylphenyl}] \mathsf{brulyl}]-2-(1,1-\mathsf{dimethylethyl})-5-\mathsf{methylphenoxy}[\mathsf{arabonyl}] \mathsf{oxy}]-5-(1,1-\mathsf{dimethylethyl})-2-\mathsf{methylphenyl}]-1-\mathsf{methylpropyl}]-2-(1,1-\mathsf{dimethylethyl})-2-\mathsf{methylphenyl}]-1-\mathsf{methylpropyl}]-2-(1,1-\mathsf{dimethylethyl})-2-\mathsf{methylphenyl}]-1-\mathsf{methylpropyl}]-2-(1,1-\mathsf{dimethylethyl})-2-\mathsf{methylphenyl}]-1-\mathsf{methylpropyl}]-2-(1,1-\mathsf{dimethylethyl})-1-\mathsf{methylphenyl}]-1-\mathsf{methylphenyl}]-1-\mathsf{methylphenylphenyl}]-1-\mathsf{methylphenylphe$

 $\label{eq:dimethylethyl} $$ \dim thyle thyl phenyl $4-3-[4-[[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methyl phenyl] butyl]-2-(1,1-dimethylethyl)-5- methyl phenoxy [carbonyl] oxyl-5-(1,1-dimethylethyl)-2-methyl phenyl]-1-[5-(1,1-dimethylethyl)-4-hydroxy-2-methyl phenyl] butyl]-2-(1,1-dimethylethyl)-5-methyl phenyl ester (9CI) (CA INDEX NAME)$

PAGE 1-A

PAGE 2-C

- L45 ANSWER 46 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN 62605-86-1
- - RL: USES (Uses)
 - (antioxidants, for thermoplastic resins)
- RN 62605-86-1 CAPLUS
 - Carbonic acid, 4-[3,3-bis[4-[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-1-methylpropyl]-2-(1,1dimethylethyl) -5-methylphenyl 4-[3-[4-[[4-[1,3-bis[5-(1,1-dimethylethyl)-

4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-1-[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenyl ester (9CI) (CA INDEX NAME)

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L45 ANSWER 47 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 63728-13-2 RL: USES (Uses)

(coatings, on photog. film supports, temperature adjustment in control of)

RN 63728-13-2 CAPLUS

Nonanedioic acid, polymer with 4,4'-(1,5-pentanediyl)bis[phenol] and 1,5-pentanediyldi-4,1-phenylene di-2-propenoate (9CI) (CA INDEX NAME)

CM 1

CRN 61469-14-5 CMF C23 H24 O4

CM

CRN 10365-62-5 CMF C17 H20 O2

CM

CRN 123-99-9 CMF C9 H16 O4

HO2C- (CH2)7-CO2H

L45 ANSWER 48 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 62605-86-1

CN

RL: MOA (Modifier or additive use); USES (Uses)
(heat stabilizers, containing phosphites, for polymers)

RN 62605-86-1 CAPLUS

62605-86-1 CAPLUS

Carbonic acid, 4-[3,3-bis[4-[[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-12-(1,1-dimethylethyl)-5-methylphenyl]-12-(1,1-dimethylethyl)-5-methylphenyl]-12-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]oxpoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-5-methylphenoxy]carbonyl]oxy]-5-(1,1-dimethylethyl)-2-methylphenyl]-1-[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]-12-(1,1-dimethylethyl)-5-methylphenyl]oxpoxy]carbonyloxy]-6(3 NIOEX NAME)

PAGE 2-C

- L45 ANSWER 49 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN IT 62605-86-1
- RL: USES (Uses)

(antioxidants, for polymers)

- RN 62605-86-1 CAPLUS
- CN Carbonic acid, 4-[3,3-bis[4-[[[4-[1,3-bis[5-(1,1-dimethylethyl)-4-hydroxy2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenoxy]carbonyl]oxy]5-(1,1-dimethylethyl)-2-methylphenyl]-1-methylpropyl]-2-(1,1dimethylethyl)-5-methylphenyl 4-[3-[4-[[[4-[1,3-bis[5-(1,1-dimethylethyl)-

4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenoxy|carbonyl]oxy|-5-(1,1-dimethylethyl)-2-methylphenyl]-1-[5-(1,1-dimethylethyl)-4-hydroxy-2-methylphenyl]butyl]-2-(1,1-dimethylethyl)-5-methylphenyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 2-B

PAGE 2-C

L45 ANSWER 50 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 62716-34-1P

RL: FORM (Formation, nonpreparative); PREP (Preparation) (formation of, in hydrolysis of secoisolariciresinol diester)

RN 62716-34-1 CAPLUS

CN 1,4-Butanedio1, 2,3-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]-, (R*,R*)(9CI) (CA INDEX NAME)

Relative stereochemistry.

L45 ANSWER 51 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 61241-77-8P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 61241-77-8 CAPLUS

CN 26,27-Dinorcholest-5-ene-3,20,22,23,24-pentol, 24-[(3\beta,17\beta)-3-(acetyloxy)androst-5-en-17-yl]-, (3\beta,20\beta)- (9CI) (CA INDEX NAME)

- L45 ANSWER 52 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- IT 52625-29-3P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 52625-29-3 CAPLUS

NN 5022-3-3 Child Co. 20 Child Co. 20 Child Co. 21 Child Ch

L45 ANSWER 53 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN ΙT

43116-70-7

RL: BIOL (Biological study) (ATPase inhibition by)

43116-70-7 CAPLUS RN

CN 1-Propanone, 1-[2-(acetyloxy)-6-(β-D-glucopyranosyloxy)-4methoxyphenyl]-3-[4-(acetyloxy)phenyl]- (CA INDEX NAME)

Absolute stereochemistry.

- L45 ANSWER 54 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- 97085-10-4P, [21,21'-Bi-17 α -pregn-4-en-20-yne]-3 β ,3' β , 17,17'-tetrol, 3,3'-diacetate RL: PREP (Preparation) ΙT
- (preparation of) RN
- 97085-10-4 CAPLUS [21,21'-Bi-17α-pregn-4-en-20-yne]-3β,3'β,17,17'-tetrol, CN
 - 3,3'-diacetate (7CI) (CA INDEX NAME)

L45 ANSWER 55 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

IT 860544-27-0P, 1,5-Pentanediol, 1,5-bis(p-hydroxyphenyl)-2,4diphenyl-, p,p'-diacetate
RL: PREP (Preparation)
 (preparation of)

RN 860544-27-0 CAPLUS

CN 1,5-Pentanediol, 1,5-bis(p-hydroxyphenyl)-2,4-diphenyl-, p,p'-diacetate (4CI) (CA INDEX NAME)

=> d bib abs 50

L45 ANSWER 50 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN

AN 1977:171037 CAPLUS

DN 86:171037

OREF 86:26853a

TI Structure of a secoisolariciresinol diester from Salvia plebeia seed

AU Powell, Richard G.; Plattner, Ronald D.

CS NRRC, ARS, Peoria, IL, USA

SO Phytochemistry (Elsevier) (1976), 15(12), 1963-5

CODEN: PYTCAS; ISSN: 0031-9422

DT Journal

LA English

01

L34 L35

AR The secoisolariciresinol diester I was isolated from S. plebeia and its structure determined on the basis of its IR, NMR, and mass spectra and those of its acetylation and hydrolysis derivs. => d his (FILE 'HOME' ENTERED AT 14:28:12 ON 12 JUN 2008) FILE 'HCAPLUS' ENTERED AT 14:29:08 ON 12 JUN 2008 SET AUTOSEARCH ON 1 US20050249857/PN FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 FILE 'HCAPLUS' ENTERED AT 14:29:29 ON 12 JUN 2008 L2 TRA L1 1- RN : 16 TERMS FILE 'REGISTRY' ENTERED AT 14:29:29 ON 12 JUN 2008 L3 16 SEA L2 10 L3 AND >=2 46.150.18/RID L4 L5 1 L3 AND OC4-C6-C6/ES 1 L3 AND C6-C6/ES 1.6 STR L8 29 L7 L9 SCR 1839 L10 29 L7 AND L9 L11 STR L7 L12 336455 >= 2 46.150.18/RID AND (OC4 OR OC4-OC4)/ES L13 50 L11 SAM SUB=L12 L14 STR L11 L15 32 L14 SAM SUB=L12 E FURAN/CN L16 1 E3 E THF/CN L17 1 E3 L18 130953 L12 AND 16.138.1/RID L19 49 L14 SAM SUB=L18 L20 STR L14 FILE 'STNGUIDE' ENTERED AT 14:55:22 ON 12 JUN 2008 FILE 'REGISTRY' ENTERED AT 14:58:52 ON 12 JUN 2008 STR L20 L22 SCR 1707 OR 1708 L23 SCR 1840 L24 2 L22 AND L23 AND L21 L25 36 L22 AND L23 AND L21 SAM SUB=L18 L26 STR L21 L27 4 L26 AND L22 AND L23 SAM SUB-L18 L28 94 L26 AND L22 AND L23 FULL SUB=L18 SAV TEM G761C1/A L28 L29 STR L7 L30 2 L29 T.31 STR L29 L32 0 L31 1.33 SCR 1701 OR 1702 OR 1703 OR 1704 OR 1705 0 L31 AND L33

33 L31 AND L33 FULL

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     FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
L37
               STR
             46 L37
L38
       694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES
L39
            50 L37 SAM SUB=L39
L40
L41
               STR L37
L42
            50 L41 SAM SUB=L39
          7953 L41 FULL SUB=L39
L43
               SAV TEM G761C1N2/A L43
    FILE 'CAPLUS' ENTERED AT 15:24:14 ON 12 JUN 2008
L44
            86 S L28
    FILE 'STNGUIDE' ENTERED AT 15:24:54 ON 12 JUN 2008
    FILE 'CAPLUS' ENTERED AT 15:30:02 ON 12 JUN 2008
    FILE 'STNGUIDE' ENTERED AT 15:31:06 ON 12 JUN 2008
    FILE 'CAPLUS' ENTERED AT 15:37:41 ON 12 JUN 2008
    FILE 'STNGUIDE' ENTERED AT 15:37:42 ON 12 JUN 2008
    FILE 'CAPLUS' ENTERED AT 15:37:57 ON 12 JUN 2008
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L45
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=> s 135
L46
           55 L35
=> d bib abs hitstr 50
L46 ANSWER 50 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
AN 1977:171037 CAPLUS
DN 86:171037
OREF 86:26853a
TI Structure of a secoisolariciresinol diester from Salvia plebeia seed
AU Powell, Richard G.; Plattner, Ronald D.
CS NRRC, ARS, Peoria, IL, USA
SO Phytochemistry (Elsevier) (1976), 15(12), 1963-5
    CODEN: PYTCAS; ISSN: 0031-9422
DT
    Journal
LA
    English
GI
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- The secoisolariciresinol diester I was isolated from S. plebeia and its AB structure determined on the basis of its IR, NMR, and mass spectra and those of its acetylation and hydrolysis derivs.
- 62716-34-1P
 - RL: FORM (Formation, nonpreparative); PREP (Preparation) (formation of, in hydrolysis of secoisolariciresinol diester)
- 62716-34-1 CAPLUS
- CN 1,4-Butanediol, 2,3-bis[[4-(acetyloxy)-3-methoxyphenyl]methyl]-, (R*,R*)-(9CI) (CA INDEX NAME)

Relative stereochemistry.

=> d bib abs hitstr 40

- L46 ANSWER 40 OF 55 CAPLUS COPYRIGHT 2008 ACS on STN
- 1982:162321 CAPLUS AN DN 96:162321
- OREF 96:26710h,26711a
- ΤI 2,3-Bis(hydroxybenzyl) derivatives
- Groen, Marinus Bernard
- AKZO N. V. , Neth. Eur. Pat. Appl., 16 pp. PA SO
- CODEN: EPXXDW
- DT Patent
- LA English EAN ONT 1

| E MIN . | CIVI | 7 | | | | | | | | | | | | | |
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| | PATENT NO. | | | | | KIND | | DATE | | | APPLICATION NO. | | | | DATE |
| | | | | | | | | | | | | | | | |
| PΙ | EP | 4315 | 0 | | | A1 | | 19820 |)106 | | EΡ | 1981-200 | 622 | | 19810605 |
| | | R: | AT, | BE, | CH, | DE, | FR. | GB, | IT, | LU, | N) | L, SE | | | |
| | ZA | 8103 | 951 | | | A | | 19820 | 0630 | | ZA | 1981-395 | 1 | | 19810611 |
| | US | 4343 | 796 | | | A | | 19820 | 0180 | | US | 1981-272 | 727 | | 19810611 |

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| | FI 8101967 | A | 19811225 | FI | 1981-1967 | 19810623 |
| | JP 57032239 | A | 19820220 | JP : | 1981-97336 | 19810623 |
| | ES 503338 | A1 | 19821101 | ES | 1981-503338 | 19810623 |
| PRAI | GB 1980-20688 | A | 19800624 | | | |
| OS | MARPAT 96:162321 | | | | | |

CH₂CH (CH₂OR²) CH (CH₂OR³) CH₂

- AB 1,4-Butanediylbis(phenols) and derivs. I [R and R1 (same or different) are OH, etherified OH, esterified OH; R2 and R3 (same or different) are H, acyl, or R2R3 = alkylidene], useful as antiinflammatory agents (no data), were prepared (±)-trans-3,4-Bis(3-hydroxybenzyl)-4,5-dihydro-2(3H)-furanone was treated with LiAlH4 in THF to give (±)-I [R = R1 = OH, R2 = R3 = H).
- IT 81436-91-1P

RL: SPN (Synthetic preparation); PREP (Preparation)

- (preparation of)
- RN 81436-91-1 CAPLUS
- CN 1,4-Butanediol, 2,3-bis[[3-(benzoyloxy)phenyl]methyl]-, (R*,R*)- (9CI) (CA INDEX NAME)

Relative stereochemistry.

| COST IN U.S. DOLLARS | SINCE FILE | TOTAL |
|--|----------------|-----------------|
| | ENTRY | SESSION |
| FULL ESTIMATED COST | 157.35 | 1075.57 |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE | TOTAL |
| CA SUBSCRIBER PRICE | ENTRY
-2 40 | SESSION
-4 8 |

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FILE CONTAINS CURRENT INFORMATION.
LAST RELOADED: Jun 6, 2008 (20080606/UP).

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L34
             0 L31 AND L33
L35
             33 L31 AND L33 FULL
                SAV TEM G761C1N/A L35
1.36
             0 L28, L35 AND L3
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FILE 'STNGUIDE' ENTERED AT 15:14:39 ON 12 JUN 2008

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FILE 'REGISTRY' ENTERED AT 15:15:43 ON 12 JUN 2008
1.37
                STR
             46 L37
L38
L39
         694251 46.150.18/RID AND (C5-C6-C6 OR C6-C6)/ES
L40
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                STR L37
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             50 L41 SAM SUB=L39
L43
           7953 L41 FULL SUB=L39
                SAV TEM G761C1N2/A L43
     FILE 'CAPLUS' ENTERED AT 15:24:14 ON 12 JUN 2008
T. 44
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     FILE 'STNGUIDE' ENTERED AT 15:24:54 ON 12 JUN 2008
     FILE 'CAPLUS' ENTERED AT 15:30:02 ON 12 JUN 2008
     FILE 'STNGUIDE' ENTERED AT 15:31:06 ON 12 JUN 2008
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    FILE 'CAPLUS' ENTERED AT 15:38:32 ON 12 JUN 2008
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            55 S L35
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     FILE 'STNGUIDE' ENTERED AT 15:43:28 ON 12 JUN 2008
=> d stat que
L31
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 8
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DEFAULT ECLEVEL IS LIMITED
GRAPH ATTRIBUTES:
RING(S) ARE ISOLATED OR EMBEDDED
NUMBER OF NODES IS 9
STEREO ATTRIBUTES: NONE
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=> log h COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION